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(54) TREATMENT OF GENOTYPED DIABETIC PATIENTS WITH DPP-IV INHIBITORS SUCH AS LINAGLIPTIN

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(58) Field of Classification Search

None

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(56) References Cited

U.S. PATENT DOCUMENTS

2,056,046 A 9/1936 Fourneau 2,375,138 A 5/1945 Victors

2,629,736	Α	2/1953	Krimmel
2,730,544	Α	1/1956	Melville
2,750,387	Α	6/1956	Krimmel
2,928,833	A	3/1960	Leake et al.
3,174,901	Α	3/1965	Sterne
3,236,891	A	2/1966	Seemuller
3,454,635	A	7/1969	Muth
3,673,241	A	6/1972	Marxer
3,925,357	A	12/1975	Okada et al.
4,005,208	A	1/1977	Bender et al.
4,061,753	A	12/1977	Bodor et al.
4,382,091	A	5/1983	Benjamin et al.
4,599,338	A	7/1986	Regnier et al.
4,639,436	A	1/1987	Junge et al.
4,687,777	A	8/1987	Meguro et al.
4,743,450	A	5/1988	Harris et al.
4,816,455	A	3/1989	Schickaneder et al.
4,873,330	A	10/1989	Lindholm
4,968,672	A	11/1990	Jacobson et al.
5,041,448	A	8/1991	Janssens et al.
5,051,509	A	9/1991	Nagano et al.
5,051,517	A	9/1991	Findeisen et al.
5,084,460	A	1/1992	Munson, Jr. et al.
5,130,244	A	7/1992	Nishimaki et al.
5,164,526	A	11/1992	Macher
5,219,870	A	6/1993	Kim
5,223,499	A	6/1993	Greenlee et al.
5,234,897	A	8/1993	Findeisen et al.
5,258,380	A	11/1993	Janssens et al.
5,266,555	A	11/1993	Findeisen et al.
5,273,995	A	12/1993	Roth
5,284,967	A	2/1994	Macher
5,300,298	A	4/1994	LaNoue
5,329,025	A	7/1994	Wong et al.
5,332,744	A	7/1994	Chakravarty et al.
		(Cont	tinued)
		(0011	

FOREIGN PATENT DOCUMENTS

AU 2003280680 A1 6/2004 AU 2009224546 A1 9/2009

(Continued)

OTHER PUBLICATIONS

Medline Plus, obesity, available at http://www.nlm.nih.gov/medlineplus/obesity.html—(referenced Aug. 22, 2013).* St. John Providence Health Center; Preventing Obesity; http://www.

stjohnprovidence.org/HealthInfoLib/swarticle.aspx?type=85&id=P07863 (referenced Aug. 22, 2013).*

eMedicine Health, diabetes causes, http://www.onhealth.com/diabetes_health/page3.htm#diabetes_causes (referenced Aug. 22, 2013).*

United Healthcare, diabetes, http://www.uhc.com/source4women/health_topics/diabetes/relatedinformation/

d0f0417b073bf110VgnVCM1000002f10bl0a_.htm—referenced Aug. 22, 2013.*

Florez, et al., "TCF7L2 polymorphisms and progression to diabetes in the diabetes prevention program," N. Eng. J. Med. 355:241-250 (2006).*

(Continued)

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(57) ABSTRACT

The present invention relates to methods for preventing or treating of metabolic disorders and related conditions, such as in certain patient groups.

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US 9,457,029 B2

Page 2

(56)		Referen	ces Cited		8,119,648 8,158,633			Himmelsbach et al. Hendrix et al.
	U.S.	PATENT	DOCUMENTS		8,138,033 8,178,541 8,232,281	B2	5/2012	Himmelsbach et al. Dugi et al.
_	200 642 4	2/1005	Daniel et al		8,455,435			Franz et al.
	5,389,642 A 5,399,578 A		Dorsch et al. Buhlmayer et al.		8,673,927			Dugi et al.
	5,407,929 A		Takahashi et al.		8,679,520		3/2014	Horres et al.
	,461,066 A		Gericke et al.		8,785,455			Hotter et al.
5	5,470,579 A		Bonte et al.		8,865,729			Sieger et al.
	,591,762 A		Hauel et al.		8,962,636 9,034,883			Pfrengle et al. Klein et al.
	5,594,003 A		Hauel et al.		9,149,478			Klein et al.
	5,602,127 A 5,614,519 A		Hauel et al. Hauel et al.		9,186,392			Klein et al.
	5,719,279 A		Kufner-Muhl et al.		2001/0020006	A1		Demuth et al.
	5,728,849 A		Bouchard et al.		2001/0051646			Demuth et al.
5	5,753,635 A		Buckman et al.		2002/0019411			Robl et al.
	5,830,908 A		Grunenberg et al.		2002/0137903 2002/0160047			Ellsworth et al. Hussain et al.
	5,879,708 A		Makino et al. Ahrndt et al.		2002/01610047			Kanstrup et al.
	5,958,951 A 5,965,555 A		Gebert et al.		2002/0169174			Chackalamannil et al.
	5,965,592 A		Buhlmayer et al.		2002/0198205	A1		Himmelsbach et al.
	5,011,049 A	1/2000	Whitcomb		2003/0040490			Sugiyama et al.
	5,107,302 A		Carter et al.		2003/0078269			Pearson et al.
	5,166,063 A		Villhauer		2003/0100563 2003/0105077			Edmondson et al. Kanstrup et al.
	5,200,958 B1 5,248,758 B1		Odaka et al. Klokkers et al.		2003/0103077			Washburn et al.
	5,303,661 B1		Demuth et al.		2003/0130313			Fujino et al.
	5,342,601 B1		Bantick et al.		2003/0149071			Gobbi et al.
	5,372,940 B1		Cavazza		2003/0153509			Bachovchin et al.
	5,448,323 B1		Jordan et al.		2003/0166578			Arch et al.
	5,548,481 B1		Demuth et al.		2003/0199528 2003/0224043		10/2003	Kanstrup et al. Appel et al.
	5,579,868 B1 5,727,261 B2		Asano et al. Gobbi et al.		2003/0232987			Dahanukar et al.
	5,784,195 B2		Hale et al.		2003/0236272		12/2003	
	5,821,978 B2		Chackalamannil et al.		2004/0023981	A1		Ren et al.
	5,869,947 B2		Kanstrup et al.		2004/0034014			Kanstrup et al.
	5,890,898 B2		Bachovchin et al.		2004/0037883 2004/0063725			Zhou et al. Barth et al.
	5,995,183 B2		Hamann et al.		2004/0063723			Himmelsbach et al.
	7,034,039 B2 7,060,722 B2		Oi et al. Kitajima et al.		2004/0077643			Yoshikawa et al.
	,000,722 B2 7,074,794 B2		Kitajima et al.		2004/0087587		5/2004	Himmelsbach et al.
	,074,798 B2		Yoshikawa et al.		2004/0097510			Himmelsbach et al.
	,074,923 B2	7/2006	Dahanukar et al.		2004/0116328			Yoshikawa et al.
	,109,192 B2		Hauel et al.		2004/0122048 2004/0122228		6/2004	Benjamin et al. Maier et al.
	7,179,809 B2		Eckhardt et al.		2004/0122228			Warne et al.
	7,183,280 B2 7,192,952 B2		Himmelsbach et al. Kanstrup et al.		2004/0138214			Himmelsbach et al.
	7,217,711 B2		Eckhardt et al.		2004/0138215			Eckhardt et al.
	,235,538 B2		Kanstrup et al.		2004/0152659			Matsuoka et al.
7	,247,478 B2		Eberhardt et al.		2004/0152720 2004/0166125			Hartig et al.
	7,291,642 B2		Kauffmann-Hefner et al.		2004/0171836		9/2004	Himmelsbach et al. Fujino et al.
	7,361,687 B2 7,393,847 B2		Barth et al. Eckhardt et al.		2004/0180925		9/2004	Matsuno et al.
	7,407,955 B2		Himmelsbach et al.		2004/0259903	A1	12/2004	Boehringer et al.
7	,407,995 B2		Ok et al.		2005/0020574			Hauel et al.
	,432,262 B2		Eckhardt et al.		2005/0026921			Eckhardt et al.
	7,439,370 B2		Eckhardt Eckhardt et al.		2005/0032804 2005/0065145			Cypes et al. Cao et al.
	7,470,716 B2 7,476,671 B2		Eckhardt et al.		2005/0070562			Jones et al.
	7,482,337 B2		Himmelsbach et al.	;	2005/0070594	A1	3/2005	Kauschke et al.
	,495,002 B2		Langkopf et al.		2005/0130985			Himmelsbach et al.
	,495,003 B2		Eckhardt et al.		2005/0143377			Himmelsbach et al.
	7,495,005 B2		Himmelsbach et al.		2005/0171093 2005/0187227			Eckhardt et al. Himmelsbach et al.
	7,501,426 B2 7,550,455 B2		Himmelsbach et al. Himmelsbach et al.		2005/0203095			Eckhardt et al.
	7,560,450 B2		Eckhardt et al.		2005/0234108			Himmelsbach et al.
	,566,707 B2		Eckhardt et al.		2005/0234235			Eckhardt et al.
7	,569,574 B2	8/2009	Maier et al.		2005/0239778			Konetzki et al.
7	7,579,449 B2		Eckhardt et al.		2005/0244502			Mathias et al. Hulin et al.
	7,610,153 B2		Carter, Jr. et al.		2005/0256310 2005/0261271			Feng et al.
	7,645,763 B2 7,718,666 B2		Himmelsbach et al. Boehringer et al.		2005/0261271			Eckhardt
	7,754,481 B2		Eberhardt et al.		2005/0266080			Desai et al.
	7,799,782 B2		Munson et al.		2005/0276794			Papas et al.
	,820,815 B2	10/2010	Pfrengle et al.		2006/0004074			Eckhardt et al.
	7,838,529 B2		Himmelsbach et al.		2006/0034922			Cheng et al.
	8,039,477 B2		Hendrix et al.		2006/0039974		2/2006	Akiyama et al.
	3,071,583 B2 3,106,060 B2		Himmelsbach Pfrengle et al.		2006/0047125 2006/0058323			Leonardi et al. Eckhardt et al.
8	5,100,000 B2	1/2012	i nengie et ai.		2000/0038323	ΑI	3/2000	Dekilarut et al.

US 9,457,029 B2

Page 3

(56)	Referen	nces Cited	2011/01				Himmelsbach et al.
U.S.	PATENT	DOCUMENTS	2011/01 2011/01				Klein et al. Dugi et al.
			2011/02				Friedl et al.
2006/0063787 A1 2006/0074058 A1		Yoshikawa et al. Holmes et al.	2011/02 2011/02				Dugi et al. Mark et al.
2006/0079541 A1		Langkopf et al.	2011/02	75561	A1	11/2011	Graefe-Mody et al.
2006/0094722 A1	5/2006	Yasuda et al.	2011/03 2012/00			1/2011	Dugi Kohlrausch et al.
2006/0100199 A1 2006/0106035 A1		Yoshikawa et al. Hendrix et al.	2012/00				Himmelsbach et al.
2006/0100033 A1 2006/0111372 A1		Hendrix et al.	2012/00				Himmelsbach et al.
2006/0111379 A1		Guillemont et al.	2012/00 2012/00				Banno et al. Graefe-Mody et al.
2006/0134206 A1 2006/0142310 A1		Iyer et al. Pfrengle et al.	2012/01				Schneider et al.
2006/0154866 A1	7/2006	Chu et al.	2012/01				Klein et al. Graefe-Mody et al.
2006/0159746 A1 2006/0173056 A1		Troup et al. Kitajima et al.	2012/01 2012/01				Sieger et al.
2006/01/5030 A1 2006/0205711 A1		Himmelsbach et al.	2012/01			6/2012	Pfrengle et al.
2006/0205943 A1		Dahanukar et al.	2012/01 2012/02				Klein et al. Himmelsbach et al.
2006/0247226 A1 2006/0270668 A1		Himmelsbach et al. Chew et al.	2012/02				Kohlrausch et al.
2006/0270701 A1		Kroth et al.	2012/02				Meinicke
2007/0027168 A1		Pfrengle et al.	2012/02 2012/02				Himmelsbach et al. Himmelsbach et al.
2007/0060530 A1 2007/0072803 A1		Christopher et al. Chu et al.	2012/02			11/2012	Sieger et al.
2007/0072810 A1	3/2007	Asakawa	2013/01				Kohlrausch et al. Klein et al.
2007/0088038 A1 2007/0093659 A1		Eckhardt et al. Bonfanti et al.	2013/01 2013/01				Pfrengle et al.
2007/0093039 A1 2007/0142383 A1		Eckhardt et al.	2013/01			8/2013	Dugi et al.
2007/0173452 A1		DiMarchi et al.	2013/02 2013/03				Ito et al. Johansen
2007/0185091 A1 2007/0196472 A1		Himmelsbach et al. Kiel et al.	2013/03				Klein et al.
2007/0197522 A1	8/2007	Edwards et al.	2014/02				Johansen et al.
2007/0219178 A1		Muramoto	2014/03 2015/01				Klein et al. Klein et al.
2007/0254944 A1 2007/0259900 A1		Hughes Sieger et al.	2015/01				Klein et al.
2007/0259925 A1	11/2007	Boehringer et al.					
2007/0259927 A1 2007/0281940 A1*		Suzuki et al. Dugi et al 514/248		FO	REIG	N PATE	NT DOCUMENTS
2007/0299076 A1		Piotrowski et al.	CA		1123	3437 A1	5/1982
2008/0039427 A1		Ray et al.	CA			5288 A1	5/1995
2008/0107731 A1 2008/0108816 A1		Kohlrausch et al. Zutter	CA			8656 A1	2/2002
2008/0234291 A1	9/2008	François et al.	CA CA			5730 A1 5249 A1	9/2002 3/2004
2008/0249089 A1 2008/0255159 A1		Himmelsbach et al. Himmelsbach et al.	CA		2496	5325 A1	3/2004
2008/0233133 A1 2008/0312243 A1		Eckhardt et al.	CA CA			3423 A1 3389 A1	4/2004 5/2004
2008/0318922 A1		Nakahira et al.	CA			3233 A1	6/2004
2009/0023920 A1 2009/0088408 A1		Eckhardt Meade et al.	CA			729 A1	12/2004
2009/0088569 A1	4/2009	Eckhardt et al.	CA CA			3074 A1 3050 A1	6/2005 9/2005
2009/0093457 A1 2009/0131432 A1		Himmelsbach et al. Himmelsbach et al.	CA		2556	5064 A1	9/2005
2009/0131432 A1 2009/0136596 A1		Munson et al.	CA CA			3067 A1 3446 A1	10/2005 10/2005
2009/0137801 A1		Himmelsbach et al.	CA			210 A1	10/2005
2009/0149483 A1 2009/0186086 A1	6/2009 7/2009	Nakahira et al. Shankar et al.	CA			2859 A1	11/2005
2009/0192314 A1	7/2009	Pfrengle et al.	CA CA			5294 A1 1912 A1	3/2006 6/2006
2009/0297470 A1 2009/0301105 A1	12/2009	Franz Loerting	CA		2599	9419 A1	11/2006
2009/0301103 A1 2009/0325926 A1		Himmelsbach	CA CA			019 A1 089 A1	11/2007 11/2007
2010/0033177 A1		Ochi et al.	CN	1		1105 A	8/2008
2010/0074950 A1 2010/0092551 A1	3/2010 4/2010	Sesha Nakamura et al.	DE			815 A1	8/1973
2010/0173916 A1	7/2010	Himmelsbach et al.	DE DE			3025 A1 3233 A1	7/1979 8/1998
2010/0179191 A1 2010/0183531 A1		Himmelsbach et al. Johncock et al.	DE		10109	0021 A1	9/2002
2010/0183331 A1 2010/0204250 A1		Himmelsbach et al.	DE DE			7803 A1 3243 A1	10/2002 3/2004
2010/0209506 A1	8/2010	Eisenreich	DE DE	1020		9540 A1	11/2005
2010/0310664 A1 2010/0317575 A1		Watson et al. Pinnetti et al.	DE	1020	04024	1454 A1	12/2005
2011/0009391 A1	1/2011	Braun et al.	DE DE			1221 A1 1054 A1	3/2006 5/2006
2011/0028391 A1		Holst et al.	EP	1020	0023	3032 A1	1/1981
2011/0046076 A1 2011/0065731 A1		Eickelmann et al. Dugi et al.	EP			0578 A2	7/1985 5/1087
2011/0092510 A1	4/2011	Klein et al.	EP EP			3403 A2 7608 A1	5/1987 9/1987
2011/0098240 A1	4/2011		EP		0248	3634 A2	12/1987
2011/0112069 A1 2011/0144083 A1		Himmelsbach et al. Himmelsbach et al.	EP EP			2675 A2 2282 A2	11/1989 9/1990
2011/01/1005 /11	5.2011				0.505		2.1220

US 9,457,029 B2

Page 4

(56)	Referenc	es Cited	WO	9929695 A1	6/1999
	FOREIGN PATEN	IT DOCUMENTS	WO WO	9938501 A2 9950248 A1	8/1999 10/1999
	FOREIGN PALEN	11 DOCUMENTS	wo	9956561 A1	11/1999
EP	0399285 A1	11/1990	WO	9967279 A1	12/1999
EP	0400974 A2	12/1990	WO	0069464 A1	11/2000
EP	409281 A1	1/1991	WO WO	0072799 A2 0073307 A2	12/2000 12/2000
EP EP	0412358 A1 443983 A1	2/1991 8/1991	WO	0107441 A1	2/2001
EP EP	0475482 A1	3/1992	WO	0132158 A2	5/2001
EP	0524482 A1	1/1993	WO	0140180 A2	6/2001
EP	0638567 A1	2/1995	WO WO	0147514 A1 0151919	7/2001 7/2001
EP	0657454 A1	6/1995 5/1997	WO	0151919	7/2001
EP EP	0775704 A1 0950658 A1	3/1997 10/1999	WO	0152825 A2	7/2001
EP	1054012 A1	11/2000	WO	0152852 A1	7/2001
EP	1066265 A1	1/2001	WO	0166548 A1	9/2001
EP	1310245 A1	5/2003	WO WO	0168603 0168646 A1	9/2001 9/2001
EP EP	1333033 1338595 A2	8/2003 8/2003	WO	0172290 A2	10/2001
EP	1406873 A2	4/2004	WO	0177110 A1	10/2001
EP	1500403 A1	1/2005	WO	0196301 A1	12/2001
EP	1514552 A1	3/2005	WO WO	0197808 A1 0202560 A2	12/2001 1/2002
EP EP	1523994 A1 1535906 A1	4/2005 6/2005	wo	0214271 A1	2/2002
EP	1537880 A1	6/2005	WO	0224698 A1	3/2002
EP	1557165 A1	7/2005	WO	02053516 A2	7/2002
EP	1586571 A1	10/2005	WO WO	02068420 A1 03000241 A2	9/2002 1/2003
EP EP	1743655 A1 1760076	1/2007	WO	03000241 A2	1/2003
EP EP	1700076 1829877 A1	3/2007 9/2007	WO	03002531 A2	1/2003
EP	1852108 A1	11/2007	WO	03002553 A2	1/2003
EP	1897892 A2	3/2008	WO WO	03004496 A1 03024965 A2	1/2003 3/2003
EP ES	2143443 A1	1/2010	WO	03024903 A2 03033686 A2	4/2003
ES ES	385302 A1 2256797 T3	4/1973 7/2006	WO	03034944 A1	5/2003
ES	2263057 T3	12/2006	WO	03037327 A1	5/2003
FR	2707641 A1	1/1995	WO	03053929 A1	7/2003
GB	2084580 A	4/1982	WO WO	03055881 A1 03057200 A2	7/2003 7/2003
HU HU	9003243 9902308 A2	5/1990 7/2000	WO	03059327	7/2003
JP	S374895 A	6/1962	WO	03064454 A1	8/2003
JP	770120	3/1995	WO	03074500 A2	9/2003
JP	8333339	12/1996	WO WO	03088900 A2 03094909 A2	10/2003 11/2003
JP JP	11193270 2000502684 A	7/1999 3/2000	wo	03099279 A1	12/2003
JР	2001213770 A	8/2001	WO	03099836 A1	12/2003
JP	2001292388 A	10/2001	WO	03103629 A1	12/2003
JР	2002348279 A	12/2002	WO WO	03104229 A1 03106428 A1	12/2003 12/2003
JP JP	2003286287 A 2003300977 A	10/2003 10/2003	WO	2004002924 A1	1/2004
JP	2004161749 A	6/2004	WO	2004011416 A1	2/2004
JP	2006045156 A	2/2006	WO	2004016587 A1	2/2004
JP	2010053576 A	3/2010	WO WO	2004018467 A2 2004018468 A2	3/2004 3/2004
JP JP	2010070576 A 2010524580 A	4/2010 7/2010	wo	2004018469 A1	3/2004
KR	20070111099 A	11/2007	WO	2004028524 A1	4/2004
WO	8706941 A1	11/1987	WO	2004033455 A1	4/2004
WO	9107945 A1	6/1991	WO WO	2004035575 A1 2004037169 A2	4/2004 5/2004
WO WO	9205175 A1 9219227 A2	4/1992 11/1992	wo	2004041820 A1	5/2004
WO	9402150 A1	2/1994	WO	2004043940	5/2004
WO	9403456 A1	2/1994	WO	2004046148 A1	6/2004
WO	9532178 A1	11/1995	WO WO	2004048379 A1 2004050658 A1	6/2004 6/2004
WO WO	9609045 A1 9611917 A1	3/1996 4/1996	wo	2004052362 A1	6/2004
wo	9636638 A1	11/1996	WO	2004058233 A1	7/2004
WO	9718814 A1	5/1997	WO	2004062689 A1	7/2004
WO	9723447 A1	7/1997	WO WO	2004065380 A1 2004074246 A2	8/2004 9/2004
WO WO	9723473 A1 9746526 A1	7/1997 12/1997	WO	2004074240 A2 2004081006 A1	9/2004
WO	9807725	2/1998	WO	2004081606 A1 2004082402 A1	9/2004
WO	9811893	3/1998	WO	2004096806 A1	11/2004
WO	9818770 A1	5/1998	WO	2004096811 A1	11/2004
WO	9822464 A1	5/1998	WO	2004106279 A2	12/2004
WO WO	9828007 A1 9840069 A2	7/1998 9/1998	WO WO	2004108730 A1 2004111051 A1	12/2004 12/2004
WO	9846082 A1	10/1998	WO	2005000846 A1	1/2004
WO	9856406 A1	12/1998	WO	2005000848 A1	1/2005

(56)	References Cited	WO 2008113000 A1 9/2008
	FOREIGN PATENT DOCUMENTS	WO 2008130998 A2 10/2008 WO 2008131149 A2 10/2008
	FOREIGN PAIENT DOCUMENTS	WO 2008137435 A1 11/2008
WO	2005007647 A1 1/2005	WO 2009011451 A 1/2009
WO	2005007658 A2 1/2005	WO 2009022007 A1 2/2009
WO	2005012288 A1 2/2005	WO 2009022008 A1 2/2009 WO 2009022009 A1 2/2009
WO	2005023179 A2 3/2005	WO 2009022009 A1 2/2009 WO 2009022010 A1 2/2009
WO WO	2005049022 A2 6/2005 2005051950 A1 6/2005	WO 2009024542 A2 2/2009
WO	2005058901 A1 6/2005	WO 2009063072 A2 5/2009
WO	2005061489 A1 7/2005	WO 2009099734 A1 8/2009
WO	2005063750 A1 7/2005	WO 2009111200 A1 9/2009 WO 2009112691 A2 9/2009
WO	2005082906 A1 9/2005	WO 2009112691 A2 9/2009 WO 2009121945 A2 10/2009
WO WO	2005085246 A1 9/2005 2005092870 A1 10/2005	WO 2009123992 A1 10/2009
WO	2005092870 A1 10/2005 2005092877 A1 10/2005	WO 2009147125 A1 12/2009
WO	2005095343 A1 10/2005	WO 2010015664 A1 2/2010
WO	2005095381 A1 10/2005	WO 2010018217 A2 2/2010 WO 2010029089 A2 3/2010
WO	2005097798 A 10/2005	WO 2010029089 A2 3/2010 WO 2010043688 A1 4/2010
WO WO	2005116000 A1 12/2005 2005116014 A1 12/2005	WO 2010045656 A2 4/2010
WO	2005117861 A1 12/2005	WO 2010072776 A1 7/2010
WO	2005117948 A1 12/2005	WO 2010079197 A1 7/2010
WO	2006005613 A1 1/2006	WO 2010086411 A1 8/2010 WO 2010092124 A1 8/2010
WO WO	2006027204 A1 3/2006	WO 2010092124 A1 8/2010 WO 2010092125 A1 8/2010
WO	2006029577 A1 3/2006 2006029769 A1 3/2006	WO 2010092163 A2 8/2010
wo	2006036664 A1 4/2006	WO 2010096384 A2 8/2010
WO	2006040625 A1 4/2006	WO 2010106457 A2 9/2010
WO	2006041976 A1 4/2006	WO 2010140111 A1 12/2010 WO 2010147768 A1 12/2010
WO WO	2006047248 A1 5/2006 2006048209 A1 5/2006	WO 2011011541 A1 1/2011
WO	2006048209 A1 5/2006 2006048427 A1 5/2006	WO 2011039337 A1 4/2011
WO	2006068163 A1 6/2006	WO 2011039367 A2 4/2011
WO	2006071078 A1 7/2006	WO 2011064352 A1 6/2011
WO	2006076231 A2 7/2006	WO 2011113947 A1 9/2011 WO 2011138380 A1 11/2011
WO WO	2006078593 A2 7/2006 2006083491 A2 8/2006	WO 2011138421 A1 11/2011
WO	2006116157 11/2006	WO 2011161161 A1 12/2011
WO	2006135693 A2 12/2006	WO 2011163206 A2 12/2011
WO	2006137085 A1 12/2006	WO 2012031124 A2 3/2012
WO	2007007173 A2 1/2007	WO 2012065993 A1 5/2012 WO 2012088682 A1 7/2012
WO WO	2007014886 A1 2/2007 2007014895 A2 2/2007	WO 2012089127 A1 7/2012
WO	2007017423 A1 2/2007	WO 2012106303 A1 8/2012
WO	2007033350 A1 3/2007	WO 2012120040 A1 9/2012
WO	2007035355 A2 3/2007	WO 2013098372 A1 7/2013 WO 2013103629 A1 7/2013
WO WO	2007035665 A1 3/2007 2007041053 A2 4/2007	WO 2013131967 A1 9/2013
WO	2007050485 A2 4/2007 2007050485 A2 5/2007	WO 2013171167 A1 11/2013
WO	2007071738 6/2007	WO 2013174768 A1 11/2013
WO	2007072083 A1 6/2007	WO 2013179307 A2 12/2013
WO	2007078726 A2 7/2007	
WO WO	2007093610 A1 8/2007 2007099345 A1 9/2007	OTHER PUBLICATIONS
WO	2007120702 A2 10/2007	Graefe-Mody et al. ("The Novel DPP-4 Inhibitor BI 1356 (proposed
WO	2007120936 A2 10/2007	tradename Ondero) and Metformin can be Safely Co-administered
WO	2007128721 A 11/2007	Without Dose Adjustment," Poster No. 553-P ADA Jun. 6-10, 2008, San Francisco—NPL 48 of IDS May 31, 2012.*
WO WO	2007128724 A1 11/2007 2007128761 A2 11/2007	Florez, et al., "TCF7L2 polymorphisms and progression to diabetes
wo	2007128761 A2 11/2007 2007135196 A2 11/2007	in the diabetes prevention program," N. Eng. J. Med. 355:241-250
WO	2007136151 A1 11/2007	(2006)*
WO	2007137107 A2 11/2007	Graefe-Mody et al. ("The Novel DPP-4 Inhibitor BI 1356 (proposed
WO	2007147185 A1 12/2007	tradename Ondero) and Metformin can be Safely Co-administered
WO WO	2007148185 A2 12/2007 2007149797 A2 12/2007	Without Dose Adjustment," Poster No. 553-P ADA Jun. 6-10, 2008,
wo	2008005569 A2 1/2008	San Francisco—N PL 48 of I DS May 31, 2012.* Verbitrava Soiji et al.: Chemical Abstract of Inparese Patent No.
WO	2008005576 A1 1/2008	Yoshikawa, Seiji et al.: Chemical Abstract of Japanese Patent No. WO 2003/104229 Preparation of purinone derivatives as
WO	2008017670 2/2008	dipeptidylpeptidase IV (DPP-IV) inhibitors, 2003.
WO WO	2008022267 A2 2/2008 2008055870 A1 5/2008	Zejc, Alfred, et al; "Badania Nad Piperazynowymi Pochodnymi
WO	2008055870 A1 5/2008 2008055940 A2 5/2008	Dwumetyloksantyn" Acta Polon Pharm, XXXV (1976) Nr. 4 pp.
WO	2008070692 A2 6/2008	417-421.
WO	2008081205 A1 7/2008	Zhong, Qing et al; "Glucose-dependent insulinotropic peptide
WO	2008083238 A2 7/2008	stimulates proliferation and TGF-? release from MG-63 cells,"
WO	2008087198 A1 7/2008	Peptides 24 (2003) 611-616.
WO	2008093878 A1 8/2008	Zimmer et al; Synthesis of 8-Substituted Xanthines and their
WO	2008093882 A1 8/2008	Oxidative Skeleton Rearrangement to 1-Oxo-2,4,7,9-

OTHER PUBLICATIONS

tetraazaspiro[4,5]dec-2-ene-6,8,10-triones; Euripean Journa Organic Chemistry (1999) vol. 9 pp. 2419-2428.

He, Y. L. et al., "Bioequivalence of Vildagliptin/Metformin Combination Tablets and Coadministration of Vildagliptin and Metformin as Free Combination in Healthy Subjects". J. Clinical Pharmacology, 2007, vol. 47, No. 9, Abstracts of the 36th Annual Meeting of the American College of Clinical Pharmacology, San Francisco, CA, Abstract 116, p. 1210.

He, Y. L. et al. "The influence of hepatic impariment on the pharmacokinetics f the dipeptidyl peptidase IV (DPP-4) inhibitor vildagliptin" European Journal of Clinical Pharmacology, vol. 63, No. 7, May 8, 2007, p. 677-686.

Huettner Silks et al: "BI 1356, a novel and selective xanthine based DPP-IV inhibitor, demonstrates good safety and tolerability with a wide therapeutic window" Diabetes

American Diabetes Association, US, vol. 56, No. Suppl 1, Jun. 1, 2007, p. A156.

International Search Report and Written Opinion for PCT/EP2010068349 mailed Feb. 4, 2011.

International Search Report and Written Opinion for PCT/EP2011/054169 mailed Aug. 4, 2011.

Januvia; Patient Information; 2010.

Jones, R.M. et al., "GPR119 agonists for the treatment of type 2 diabetes". Expert Opinion on Therapeutic Patents 2009 Informa Healthcare for GBR LNKSD—DOI: 10.1517/13543770903153878, vol. 19, No. 10, Oct. 2009, p. 1339-1359.

Kanada, S. et al., "Safety, tolerability, pharmacokenetics and pharmacodynamics of multiple doses of BI 1356 (proposed tradename Ondero), a dipeptidyl peptidase 4 inhibitor, in Japanese patients with type 2 diabetes" Diabetes, vol. 57, No. Suppl. 1, Jun. 2008, p. A158-A159 and 68th Annual Meeting of the American Diabetes Association: San Francisco, CA, Jun. 6-10, 2008.

 $\label{eq:Kim} Kim, \quad D. \quad \text{et} \quad al., \quad \text{``(2R)-4-Oxo-4-(3-(Trifluoremethyl)-5,6-dihydro[1,2,4]triazolo[4,3-a]pyrazin-7(8H)-yl]-1-(2,4,5-a)} \\$

trifluorophenyl)butan-2-amine: A Potent, Orally Active Dipeptidyl Peptidase IV inhibitor for the Treatment of Type 2 Diabetes." Journal Med. Chem, 2005, 48, p. 141-151.

Knorr, M. et al., "Comparison of Direct and Indirect Antioxidant Effects of Linagliptin (BI 1356, Ondero) with other Gliptins—Evidence for Anti-Inflammatory Properties of Linagliptin". Free Radical Biology and medicine, Elsevier Science, U.S. vol. 49, Oct. 23, 2010, p. S197.

Korom, S. et al; Inhibition of CD26/dipeptidyl peptidase IV activity in vivo prolongs cardiac allograft survival in rat recipients1,2, Transplantation, May 27, 1997, vol. 63, No. 10, pp. 1495-1500.

Lambier, A.M. et al., Dipeptidyl-Peptidase IV from Bench to Bedside: An Update on Structural Properties, Functions, and Clinical Aspects of the Enzyme DPP IV. Critical Reviews in Clinical Laboratory Sciences, 2003, 40(3), p. 209-294.

Levien, T.L. et al, "New drugs in development for the treatment of diabetes", Diabetes Spectrum, American Diabetes Association, US, vol. 22, No. 2, 1 Jan. 2009, pp. 92-106.

March, J. "Advanced Organic Chemistry: Reactions, Mechanisms, and Structure". Fourth Edition, 1992, pp. 652-653.

Meece, J. "When Oral Agents Fail: Optimizing Insulin Therapy in the Older Adult". Consultant Pharmacist, The Society, Arlington, VA US. vol. 24, No. Suppl B, Jun. 1, 2009, p. 11-17.

Mendes, F.D, et al. "Recent advances in the treatment of non-alcoholic fatty liver disease". Expert Opinion on Investigational Drugs, vol. 14, No. 1, Jan. 1, 2005, p. 29-35.

Merck: "Initial Therapy with Janumet (sitagliptin/metformin) provided significantly greater blood sugar lowering compared to metformin alone in patients with type 2 diabetes". Webwire.com, Jun. 8, 2009, p. 1-4. http://www.webwire.com/ViewPressRel.asp?ald=96695.

O'Farrell, et al., "Pharmacokinetic and Pharmacodynamic Assessments of the Dipeptidyl Peptidase-4 Inhibitor PHX1149: Double-Blind, Placebo-controlled, Single-and Multiple-Dose Studies in Healthy Subjects". Clinical Therapeutics, Excerpta Medica, Princeton, NJ, vol. 29, No. 8, 2007, p. 1692-1705.

Patani George A. et al.: "Bioisoterism: A Rational Approach in Drug Design", Chemical Reviews, 1996, vol. 96, No. 8, pp. 3147-3176.

Pei, Z.: "From the bench to the bedside: Dipeptidyl peptidase IV inhibitors, a new class of oral antihyperglycemic agents" Current Opinion in Drug Discovery and Development, Current Drugs, London, GB vol. 11, No. 4, Jul. 1, 2008 pp. 512-532.

Pospisilik, et al; Dipeptidyl Peptidase IV Inhibitor Treatment Stimulates? -Cell Survival and Islet Neogenesis in Streptozotocin-Induced Diabetic Rats; Diabetes, vol. 52, Mar. 2003 pp. 741-750. Priimenko, B. A., et al; Synthesis and Pharmacological Activity of Derivates of 6,8-Dimethyl Imidazo(1,2-f) Xanthine-(Russ.); Khimiko-Farmatsevticheskii zhurnal (1984) vol. 18, No. 12 pp. 1456-1461.

Rhee et al.: "Nitrogen-15-Labeled Deoxynucleosides. 3. Synthesis of [3-15N]-2'-Deoxyadenosine" J. Am. Chem. Soc. 1990, 112, 8174-8175.

Rosenstock, et al., "Efficacy and tolerability of initial combination therapy with vildagliptin and pioglitazone compared with component montherapy in patients with type 2 diabetes". Diabetes, Obesity and Metabolism, Mar. 2007, vol. 9, No. 2, p. 175-185.

Rosenstock, J. et al., "Alogliptin added to insulin therapy in patients with type 2 diabetes reduces HbA1c without causing weight gain or increased hypoglycaemia". Diabetes, Obesity and Metabolishm, Dec. 2009, vol. 11. No. 12, p. 1145-1152.

Salomon, J., et al; Ultraviolet and g-Ray-Induced Reactions of Nucleic Acid Constituents. Reactions of Purines with Amines; Photochemistry and Photobiology (1974) vol. 19 pp. 21-27.

Sathananthan, A., et al., "Personalized pharmacotherapy for type 2 diabetes mellitus". Personalized Medicine 2009 Future Medicine Ltd, vol. 6, No. 4, Jul. 2009, p. 417-422.

Sauer, R, et al. "Water-soluble phosphate prodrugs of 1-Propargyl-7-styrylxanthine derivatives, A2A-selective adenosine receptor antagonists". Journal Med. Chem., vol. 43, Issue 3, Jan. 2000, p. 440-448.

Schwartz, M. S. et al., "Type 2 Diabetes Mellitus in Childhood: Obesity and Insulin Resistance"—JAOA Review Article, vol. 108, No. 9, Sep. 2008, p. 518.

Scientific Discussion: "Eucreas. Scientific discussion". Online Oct. 2007, p. 1-27, URL:http://www.emea.europa.eu/humandocs/PDFs/EPAR/eucreas/H-807-en6.pdf. see point 2. quality aspects pp. 2-4. (EMEA).

Sedo, A. et al; "Dipeptidyl peptidase IV activity and/or structure homologs: Contributing factors in the pathogenesis of rheumatoid arthritis?" Arthritis Research & Therapy 2005, vol. 7, pp. 253-269. Stahl, P.H., "Handbook of Pharmaceutical Salts". C.G. Wermuth, Wiley-VCH, 2002, p. 61.

Sune Negre, J. M. "New Galenic Contributions to Administration Forms", Continued Training for Hospital Pharmacists 3.2. [retrieved on Feb. 23, 2011]. Retrieved from the internet http://www.ub.es/legmh/capitols/sunyenegre.pdf>.

Tamm, E, et al., "Double-blind study comparing the immunogenicity of a licensed DTwPHib-CRM197 conjugate vaccine (Quattvaxem TM) with three investigational, liquuid formulations using lower doses of Hib/CRM197 conjugate". Science Direct, Vaccine, Feb. 2005, vol. 23, No. 14, p. 1715-1719.

Tanaka, S.. et al; "Suppression of Arthritis by the Inhibitors of Dipeptidyl Peptidase IV," In. J. Immunopharmac., vol. 19, No. 1, pp. 15-24, 1997.

Thomas, L, et al: "BI 1356, a novel and selective xanthine beased DPP-IV inhibitor, exhibits a superior profile when compared to sitagliptin and vildagliptin." Diabetologoa, vol. 50, No. Suppl. 1, Sep. 2007, p. S363.

Thomas, L., "Chronic treatment with the Dipeptidyl Peptidase-4 Inhibitor BI 1356[9R)-8-(3-Amino-piperidin-1-yl)-7-but-2-ynyl-3-methyl-1(4-methyl-quinazolin-2-ylmethyl)-3,7-dihydro-purine-2,6-dione] Increases Basal Glucagon-Like Peptide-1 and Improves Glycemic Control in Diabetic Rodent Models" The Journal of Pharmacology and Experimental Therapeutics, Feb. 2009, vol. 328, No. 2, pp. 556-563.

Thomas, Leo et al: "(R)-8-(3-Amino-piperidin-1-yl)-7-but-2-ynyl-3-methyl-1-(4-methyl-quinazolin-2-ylmethyl)-3,7-dihydro-purine-2,6-dione (BI 1356), a Novel Xanthine-Based Dipeptidyl Peptidase

OTHER PUBLICATIONS

4 Inhibitor, Has a Superior Potency and Longer Duration of Action Compared with Other Dipeptidyl Peptidase-4 Inhibitors" Journal of Pharmacology and Experimental Therapeutics, American Socity for Therapeutics, US, vol. 325, No. 1, Apr. 1, 2008, pp. 175-182.

U.S. Appl. No. 12/724,653, filed Mar. 16, 2010—Xanthine Derivatives, the Preparation Thereof and Their Use as Pharmaceutical Compositions. Inventor: Frank Himmelsbach, et al.

U.S. Appl. No. 12/767,855, filed Apr. 27, 2010—Xanthine Derivatives, the Preparation Thereof and Their use as Pharmaceutical Compositions. Inventor: Frank Himmelsbach, et al.

Villhauer, E.B., "1-[[3-Hydroxy-1-adamantyl)amino]acetyl]-1-cyano-(S)-pyrrolidine: A Potent, Selective, and Orally Bioavailable Dipeptidyl Peptidase IV Inhibitor with Antihyperglycemic Properties" Journal Med. Chem, 2003, 46, p. 2774-2789.

Villhauer, E.B., et al., "1-{2-{5-Cyanopyridin-2-yl)amino}-ethylamino}acetyl-1-1(S)-pyrrolidine-carbonitrile: A Potent, Selective, and Orally Bioavailable Dipeptidyl Peptidase IV Inhibitor with Antihyperglycemic Properties". Journal of Medical Chemistry, 2002, vol. 45, No. 12, p. 2362-2365.

Wang Y et al: "BI-1356. Dipeptidyl-peptidase IV inhibitor, antidiabetic agent" Drugs of the Future, Prous Science, ES,vol. 33, No. 6, Jun. 1, 2008, pp. 473-477.

White, J.R., "Dipeptidyl Peptidase-IV Inhibitors: Phamacological Profile and Clinical Use". Clinical Diabetes, vol. 26, 2008, p. 53-57. Wikipedia, Annulation. Jun. 23, 2008, http://en.wikipedia.org/wiki/Annelation

Williams-Herman, D. et al., "Efficacy and safety of initial combination therapy with sitagliptin and metformin in patients with type 2 diabetes: a 54-week study". Current Medical Research and Opinion, Informa Healthcare, GB, vol. 25, No. 3, Jan. 2009, p. 569-583.

Wolff, M.E.: "Burger's Medicinal Chemistry and Drug Discovery" Fifth Edition, vol. 1: Principles and Practice, pp. 975-977, 1994, John Wiley & Sons, Inc.

World Health Organization (WHO). "Addendum 1 to "The use of stems in the selection of International Nonproprietary names (INN) for pharmaceutical substances" Online Jun. 19, 2007, pp. 1-3, retrieved from URL: http://www.who.int/medicindedocs/index/assoc/s1414e/s1414e.pdf.

X-Ray Diffraction. The United States Pharmacopeia, 2002, USP 25 NF20, p. 2088-2089.

Yasuda, et al. "E3024 3-but-2-ynyl-5-methyl-2-piperazin-1-yl-3,5-dihydro-4H-imidazol [4,5-d]pyridazin-4-one tosylate, is a move, selective and competitive dipeptidyl peptidase-IV inhibitor". European Journal of Pharmacology, vol. 548, No. 1-3, Oct. 24, 2006, p. 181-187. Abstract.

Abstract in English for German DE2205815, 1972.

Abstract in English for German EP0023032, 1981.

Abstract in English, for KR20070111099, Nov. 11, 2007.

Ahren B: "DPP-4 inhibitors", Best practice and research in clinical endocrinology and metabolism—New therapies for diabetes 200712 GB LNKD—DOI:10.1016/J. Beem.2007.07.005, vol. 21, No. 4, Dec. 2007, pp. 517-533.

Augeri, D.J. "Discovery and Preclinical Profile of Saxagliptin (GMB-477118): A Highly Potent, Long-Acting, Orally Active Dipeptidyl Peptidase IV Inhibitor for the Treatment of Type 2 Diabetes". Journal Med. Chem, 2005, vol. 48, No. 15, p. 5025-5037. Augusti, D.V. et al., "Quantitative determinatio of the enantiomeric composition of thalidomide solutions by electrospray ionizatio tandem mass spectrometry". Chem Comm, 2002, p. 2242-2243.

Augustyns, K. et al., The Unique Properties of Dipeptidyl-peptidase IV (DPP IV/CD 26) and the Therapeutic Potential of DPP-IV Inhibitors, Current Medicinal Chemistry, vol. 6, No. 4, 1999, pp. 311-327.

Aulinger, B.A. et al., "Ex-4 and the DPP-IV Inhibitor Vildagliptin have Additive Effects to Suppress Food Intake in Rodents". Abstract No. 1545-P. 2008.

Balaban, Y.H.et al., "Dipeptidyl peptidase IV (DDP IV) in NASH patients" Annals of Hepatology, vol. 6, No. 4, Oct. 1, 2007, pp. 242-250, abstract.

Balkan, B. et al, "Inhibition of dipeptidyl peptidase IV with NVP-DPP728 increases plasma GLP-1 (7-36 amide) concentrations and improves oral glucose tolerance in obses Zucker rates". Diabetologia, 1999, 42, p. 1324-1331.

Bastin, R.J. et al., "Salt Selection and Optimization Procedures for Pharmaceutical New Chemical Entities". Organic Process Research and Development, 2000, vol. 4, p. 427-435.

Beljean-Leymarie et al., Hydrazines et hydrazones hétérocycliques. IV. Synthèses de dérivés de l'hydrazine dans la série des imidazo[4,5-d]pyridazinones-4, Can. J. Chem., vol. 61, No. 11, 1983, pp. 2563-2566.

Bollag, R.J. et al; "Osteoblast-Derived Cells Express Functional Glucose-Dependent Insulinotropic Peptide Receptors," Endocrinology, vol. 141, No. 3, 2000, pp. 1228-1235.

Brazg, R. et al: "Effect of adding sitagliptin, a dipeptidyll peptidase-4 inhibitor, to metformin on 24-h glycaemic control and beta-cell function in patients with type 2 diabetes." Diabetes, Obesity and Metabolism, Mar. 2007, vol. 9, No. 2, Mar. 2007 pp. 186-193.

Brittain, H.G., "Methods for the Characterization of Polymorphs: X-Ray Powder Diffraction," Polymorphism in Pharmaceutical Solids, 1999, p. 235-238.

Bundgaard, H. "Design of prodrugs: Bioreversible derivatives for various functional groups and chemical entities". Royal Danish School of Pharmacy, 1985, p. 1-92.

Busso et al., "Circulating CD26 is Negatively Associated with Inflammation in Human and Experimental Arthritis," Am. J. Path., vol. 166, No. 2, Feb. 2005, pp. 433-442.

Caira, M.R., "Crystalline polymorphism of organic compounds" Topics in Current Chemistry, Springer, Berlin, vol. 198, 1998, p. 163-208.

Chemical Abstract. EP412358, 1991:185517, Findeisen.

Chemical Abstract: FR2707641, 1995:543545, Dodey.

Chemical Abstracts Accession No. 106:95577 Romanenko et al., "Synthesis and Biological Activity of 3-Methyl, 7- or 8-alkyl-7,8dialkyl, heterocyclic, and cyclohexylaminoxanthines," Zaporozh. Med. Institute (1986).

Chemical Abstracts Accession No. 1987:95577: Abstract of Romanenko et al., "Synthesis and biological activity of 3-methyl, 7-or 8-alkyl, 7,8-dialkyl, heterocyclic, and cyclohexylaminoxanthines," Zapoeozh, USSR, Farmatsevtichnii Zhurnal, 1986, (Kiev), vol. 5, 1986, pp. 41-44.

Clinical Trials. "View of NCT00601250 on Jan. 25, 2008: Efficacy and Safety of BI 1356 vs Placebo added to Metformin Background Therapy in Patients with Type 2 Diabetes" Clinical Trials. Gov Archive, [Online] Jan. 25, 2008 URL:http://clinicaltrials.gov/archive/NCT00601250/2008_01_25 [retrieved on Feb. 27, 2009].

Clinical Trials. NCTO0622284. "Efficacy and safety of BI 1356 in combination with metformin in patients with type 2 diabetes" ClinicalTrials.gov (Online) No. NCT00622284, Feb. 13, 2008, p. 1-5, URL:http://clinicaltrial.gov/ct2/show/.

Clinical Trials. View of NCT00730275 updated on Aug. 7, 2008. "A study to assess the pharmacokinetics, safety and tolerability of Sitagliptin in adolescents". http://clinicaltrials.gov/archive/NCT00730275/2008_08_07.

Clinical Trials: NCT00954447, View on Jun. 14, 2010. "Efficacy and Safety of Linagliptin in Combination with Insulin in Patients with Type 2 Diabetes". http://clinicaltrials.gov/archive/NCT0095444712010_06_14.

Clinical Trials: NCT00309608. Efficacy and safety of BI 1356 in combination with metformin in patients with type2 diabetes. Boehringer Ingelheim Pharmaceuticals, Jan. 27, 2009. Clinical Trials.gov . http://clinicaltrials.gov/archive/NCT00309608/2009_01_27.

Clinical Trials: NCT00602472. "BI 1356 in combination withe metformin and a sulphonylurea in Type 2 Diabetes". DrugLib.com, Nov. 3, 2008. http://www.druglib.com/trial/08/NCT00309608.html. Clinical Trials: NCT00622284. Efficacy and Safety of BI 1356 in Combination with Metformin in Patients with Type 2 Diabetes.

OTHER PUBLICATIONS

Boehringer Ingelheim Pharmaceuticals, Aug. 2008. http://clinicaltrials.gov/archive/NCT00622284/2010_01_13.

Clinical Trials: NCT00798161. "Safety and efficacy of Bi 1356 Plus Metformin in Type 2 Diabetes, Factorial Design". Clinical Trials. gov archive. A Service of the U.S> National Institutes of Health. Nov. 24, 2008, p. 1-3. http://clinicaltrials.gov/archive/NCT00798161/2008_11_24.

Combs, D. W. et al., "Phosphoryl Chloride Induced Ring Contraction of 11,4-Benzodiazepinones to Chloromethylquinazolines". J. Heterocyclic Chemistry, BD. 23, 1986, p. 1263-1264.

Conarello, S.L. et al., "Mice lacking dipeptidyl peptidase IV are protected against obesity and insulin resistance". PNAS, May 27, 2003, vol. 100, No. 11, p. 6825-6830.

Cygankiewicz, Andrzej et al., Investigations into the Piperazine Derivatives of Dimethylxanthine:, Acta Polon. Pharm. [Papers of Polish Pharmacology], XXXOV, No. 5, pp. 607-612, 1977.

Dave, K.G. et al., "Reaction of Nitriles under Acidic Conditions, Part I. A General Method of Synthesis of Condensed Pyrimidines", J. Heterocyclic Chemistry, BD, 17, 1, ISSN 0022-152X,Nov. 1980, p. 1497-1500.

Deacon, C.F. et al; "Dipeptidyl peptidase IV inhabitation as an approach to the treatment and prevention of type 2 diabetes: a historical perspective;" Biochemical and Biophysical Research Communications (BBRC) 294 (2002) 1-4.

Deacon, C.F., et al. Inhibitors of dipeptidyl peptidase IV: a novel approach for the prevention and treatment of Type 2 diabetes? Expert Opinion on Investigational Drugs, Sep. 2004, vol. 13, No. 9, p. 1091-1102.

DeMeester, I. et al.; "CD26, let it cut or cut it down", Review: Immunology Today; Aug. 1999, vol. 20, No. 8 pp. 367-375.

Dugi, K.A. et al., "BI 1356, a novel xanthine-based DPP-IV inhibitor, exhibits high potency with a wide therapeutic window and significantly reduces postprandial glucose excursions after an oGIT". Diabetologia, vol. 50, No. Suppl 1, Sep. 2007, pS367, and 43rd Annual Meeting of the European Association for the Study of Diabetes; Amsterdam, Netherlands, Sep. 18-21, 2007.

Eckhardt Matthias et al: 8-(3-(R)-aminopiperidin-1-yl)-7-but-2-yny 1-3-methyl-1-(4-methyl-quina zolin-2-ylmethyl)-3,7-dihydropurine-2,6-dione (BI 1356), a highly potent, selective, longacting, and orally bioavailable DPP-4 inhibitor nhibitor for the treatment of type 2 diabetes: Journal of Medicinal Chemistry, American Chemical Society. Washington.; US, vol. 50, No. 26, Dec. 1, 2007, p. 6450-6453.

Eckhardt, M. et al., "3,5-dihydro-imidazo[4,5-d]pyridazin-4-ones: a class of potent DPP-4 inhibitors" Bioorganic & Medicinal Chemistry Letters, Pergamon, Elsevier Science, GB, vol. 18, No. 11, Jun. 1, 2008, pp. 3158-3162, XP022711188.

Elrishi M A et al: "The dipeptidyl-peptidase-4 (D::-4) inhibitors: A new class of oral therapy for patients with type 2 diabetes mellitus" Practical Diabetes International Chichester, vol. 24, No. 9, Nov. 1, 2007 pp. 474-482.

Florez, Jose C., et al., "TCF7L2 Polymorphisms and progression to diabetes in the diabetes prevention program". New England Journal of Medicine, MA Medical Society, vol. 355, No. 2, Jul. 20, 2006, p. 241-250.

Gallwitz, B. et al., "Saxagliptin, a dipeptidyl peptidase Iv inhibitor for the treatment of type 2 diabetes". IDrugs, vol. 11, No. 12, Dec. 2008, p. 906-917.

Garber, A.J. et al., "Update: Vildaglitin for the treatment of Type 2 diabetes" Expert Opinion on Investigational Drugs, 200801GB, vol. 17, No. 1, Jan. 2008, p. 105-113.

Garcia-Soria, et al., "The dipeptidyl peptidase-4 inhibitor PHX1149 improves blood glucose control in patents with type 2 diabetes mellitus". Diabetes, Obesity and Metabolism, Apr. 2008, vol. 10, No. 4, p. 293-300.

Gennaro, Alfonso, R; Remington: The Science and Practice of Pharmacy: Oral Solid Dosage Forms; Mack Publishing Company, Philadelphia, PA (1995) vol. II, 19th Edition, Ch. 92 pp. 1615-1649. Giron, D.; Thermal Analysis and Calorimetric Methods in the Characterisation of Polymorphs and Solvates; Thermochimica Acta (1995) vol. 248 pp. 1-59.

Graefe-Mody et al., "The novel DPP-4 inhibitor . . ." Diabetes, (online) 2008, XP002561421 http://professional.diabetes.org/content/posters/2008/p553-p.pdf.

Greene, T.W, et al., "Protection for the Amino Group". Protective Groups in Organic Synthesis, 3rd edition, 1999, p. 494-653.

Gwaltney, S. "Medicinal Chemistry Approaches to the Inhibition of Dipeptidyl Peptidase IV", Current Topics in Medicinal Chemistry, 2008, 8, p. 1545-1552.

Abstract in English for German DE10109021, 2002.

Ahren, Bo, et al; Improved Meal-Related b-Cell Function and Insulin Sensitivity by the Dipeptidyl Peptidase-IV Inhibitor Vildagliptin in Metformin-Treated Patients with Type 2 Diabetes Over 1 Year; Diabetes Care (2005) vol. 28, No. 8 pp. 1936-1940. American Diabetes Association, "Standards of Medical Care in Diabetes-2008." Diabetes Care, Jan. 2008, vol. 31, Supplement 1, pp. S12-S54.

Anonymous, Clinicaltrials.gov, 2008, No. NCT00622284, "Efficacy and Safety of BI 1356 in combination with metformin in patients with type 2 diabetes" p. 1-5.

Anstee, Quentin M. et al. "Mouse models in non-alcholic fatty liver disease and steatohepatitis research" (2006) International Journal of Expermental Pathology, vol. 87, pp. 1-16.

Balbach, S. et al., "Pharmaceutical evaluation of early development candidates the 100 mg-approach." International Journal of Pharmaceutics, 2004, vol. 275, pp. 1-12.

Berge, S. et al., "Pharmaceutical Salts." Journal of Pharmaceutical Sciences, 1977, vol. 66, No. 1, pp. 1-19.

Bernstein, Joel "Polymorphism in Molecular Crystals." Oxford University Press, 2002, p. 9.

Bosi, E. et al., "Effects of Vildagliptin on Glucose Control Over 24 Weeks in Patients With Type 2 Diabetes Inadequately Controlled With Metformin." Diabetes Care, 2007, vol. 30, No. 4, pp. 890-895. Brazg, Ronald, et al; Effect of Adding MK-0431 to On-Going Metforming Therapy in Type 2 Diabetic Patients Who Have Inadequate Glycemic Control on Metformin; Diabetes ADA (2005) vol. 54, Suppl. 1 p. A3.

Byrn, Stephen R. "Solid-State Chemistry of Drugs." Academic Press, 1982, pp. 1-27.

Campbell, R. Keith "Rationale for Dipeptidyl Peptidase 4 Inhibitors: A New Class of Oral Agents for the Treatment of Type 2 Diabetes Mellitus." The Annals of Pharmacotherapy, Jan. 2007, vol. 41, pp. 51-60.

Charbonnel, B. et al., "Efficacy and Safety of the Dipeptidyl Peptidase-4 Inhibitor Sitagliptin Added to Ongoing Metformin Therapy in Patients With Type 2 Diabetes Inadequately Controlled With Metformin Alone." Diabetes Care, 2006, vol. 29, No. 12, pp. 2638-2643

Chemical Abstracts Service, Database Accession number No. RN 668270-12-01, 2004, "1H-Purine-2,6-dione, 8-[(3R)-3-amino-1-piperidinyl]-7-(2-butyn-1-yl)-3,7-dihydro-3-methyl-1-[(4-methyl-2-quinazolinyl)methyl]".

Chemistry Review: Tradjenta, "NDA 201280, CMC Director Review Tradjenta (Linagliptin) Tablets." Center for Drug Evaluation and Research, Aug. 9, 2010, Retrieved from the internet on Nov. 1, 2013, http://www.accessdata.fda.gov/drugsatfda_docs/nda/20111201280Orig1s000ChemR.pdf.

Clinical Trial NCT00622284 (published online at clinicaltrials.gov on Feb. 22, 2008).

Conarello, S.L. et al; "Mice lacking dipeptidyl peptidase IV are protected against obesity and insulin resistance," PNAS 2003; 100:6825-6830; originally published online May 14, 2003; information current as of Dec. 2006. www.pnas.org/cgi/content/full/100/11/6825.

Dave, Rutesh H. "Overview of pharmaceutical excipients used in tablets and capsules." Drug Topics, Oct. 24, 2008.

Deacon, Carolyn F., et al., "Linagliptin, a xanthine based dipeptyl peptidase-4 inhibitor with an unusual profile for the treatment of type 2 diabetes" Expert Opinion Investig. Drugs 2010, 19 (1) p. 133-140.

OTHER PUBLICATIONS

Deacon, Carolyn F. et al. "Linaglipitn, a xanthine-based dipeptidyl peptidase-4 inhibitor with an unusual profile for the treatment of type 2 diabetes" Expert Opin. Investig. Drugs (2010) 19(1): 133-140

Definition of "prevent", e-dictionary, Aug. 15, 2013, http://dictionary.reference.com/browse/prevent.

Demuth, H-U. et al., "Type 2 diabetes—Therapy with dipeptidyl peptidase IV inhibitors". Biochimica et Biophysica Acta, vol. 1751(1), 2005, p. 33-44.

Diabetes Health Center, "Diabetic Retinopathy—Prevention." Retrieved online Mar. 22, 2011. www.diabetes.webmd.com/tc/diabetic-retinopathy-prevention http://www.diabetes.webmd.com/tc/diabetic-retinopathy-prevention?print=true.

Diabetesincontrol.com "EASD: Eucreas, a Combination of Galvus and Metformin, Recommended for Approval." Diabetes in Control. com, Sep. 25, 2007, Retrieved from internet on Nov. 30, 2012, http://www.diabetesincontrol.com/articles/53-diabetes-news/5145. Diabetic Neuropathy, Retrieved online Mar. 6, 2012. www.mayoclinic.com/health/diabetic-neuropathy/DS01045/

METHOD=print&DSE http://www.mayoclinic.com/health/diabetic-neuropathy/DS01045/METHOD=print&DSE.

Drucker, et al.., The incretin system:glucagon-like peptide-1 receptor agonists and dipeptidyl peptidase-4 inhibitors in type 2 diabetes. Lancet, 2006, 368: 1696-705.

Dunitz, J. et al., "Disappearing Polymorphs." Acc. Chem. Res. 1995, vol. 28, No. 4, pp. 193-200.

Edosada, C. Y. et al. "Selective Inhibition of Fibroblast Activation Protein Protease Based on Dipeptide Substrate Specificity." The Journal of Biological Chemistry, 2006, vol. 281, No. 11, pp. 7437-7444.

eMedicine Health, "Diabetes Causes." Retrieved from internet on Aug. 22, 2013. httm#diabetes causes>.

Eucreas Scientific Discussion, 2007, p. 1-27, www.emea.europa.eu/humandocs/PD/Fs/EPAR/eucreas/H-807-en6.pdf, Anonymous.

European Search Report for EP 08 15 9141 mailed Apr. 6, 2009 (European counterpart of U.S. Appl. No. 12/143,128).

Ferreira, L. et al., "Effects of Sitagliptin Treatment on Dysmetabolism, Inflammation, and Oxidative Stress in an Animal Model of Type 2 Diabetes (ZDF Rat)." Mediators of Inflammation, 2010, vol. 2010, pp. 1-11.

Forst, T. et al., "The Novel, Potent, and Selective DPP-4 Inhibitor BI 1356 Significantly Lowers HbA1c after only 4 weeks of Treatment in Patients with Type 2 Diabetes." Diabetes, Jun. 2007, Poster No. 0594P.

Fukushima et al., Drug for Treating Type II Diabetes (6), "action-mechanism of DPP-IV inhibitor and the availability thereof" Mebio, 2009, vol. 26, No. 8, p. 50-58.

Gallwitz, B. "Sitagliptin with Metformin: Profile of a Combination for the Treatment of Type 2 Diabetes". Drugs of Today, Oct. 2007, 43(10), p. 681-689.

Gallwitz, B. et al., DPP IV inhibitors for the Treatment of Type 2 Diabetes; Diabetes Frontier (2007) vol. 18, No. 6 pp. 636-642.

Garber, A. J. et al., "Effects of Vildagliptin on Glucose Control in Patients with Type 2 Diabetes Inadequately Controlled with a Sulphonylurea". Diabetes, Obesity and Metabolism (2008) vol. 10 pp. 1047-1055.

Gennaro, Alfonso R., Remington Farmacia, 19th Edition, Spanish copy, 1995, p. 2470.

Graefe-Mody, et al; Evaluation of the Potential for Steady-State Pharmacokinetic and Pharmacodynamic Interactions Between the DPP-4 Inhibitor Linagliptin and Metformin in Healthy Subjects; Currents Medical Research and Opinion (2009) vol. 25, No. 8 pp. 1963-1972.

Halimi, et al., "Combination treatment in the management of type 2 diabetes focus on vildagliptin and metformin as a single tablet", Vascualr Health and Risk Management, 2008, 4(3) p. 481-92.

Hayashi, Michio., "Recipe for Oral Hypoglycemic Agents to Pathological Condition" Pharmacy (2006) vol. 57, No. 9 pp. 2735-2739.

Heise, T. et al., "Treatment with BI 1356, a Novel and Potent DPP-IV Inhibitor, Significantly Reduces Glucose Excursions after an oGTT in Patients with Type 2 Diabetes." A Journal of the American Diabetes Association, Jun. 2007, vol. 56, Supplement 1, Poster No. 0588P.

Herman, G. A. et al., "Dipeptidyl Peptidase-4 Inhibitors for the Treatment of Type 2 Diabetes: Focus on Sitagliptin." Clinical Pharmacology and Therapeutics, 2007, vol. 81, No. 5, pp. 761-767. Herman, Gary et al. "Co-Administration of MK-0431 and Metformin in Patients with Type 2 Diabetes Does Not Alter the Pharmacokinetics of MK-0431 or Metformin" (2005) Journal of American Diabetes Association vol. 54, Supplement 1, 3 pgs.

Hermann, Robert, et al; Lack of Association of PAX4 Gene with Type 1 Diabetes in the Hungarian Populations; Diabetes (2005) vol. 54 pp. 2816-2819.

Hermansen, K., "Efficacy and Safety of the Dipeptidyl Peptidase-4 Inhibitor, Sitagliptin, in Patients with Type 2 Diabetes Mellitus Inadequately Controlled on Glimepiride Alone or on Glimepiride and Metformin". Diabetes, Obesity and Metabolism (2007) vol. 9, No. 5 pp. 733-745.

Hilfiker, R. et al., "Relevance of Solid-state Properties for Pharmaceutical Products." Polymorphism in the Pharmaceutical Industry, 2006, Chapter 1, pp. 1-19.

Horsford, E. N. "On the source of free hydrochloric acid in the gastric juice." Proceedings of the Royal Society of London, Published in 1868-1869, vol. 17, pp. 391-395.

Hu, Y. et al., "Synthesis and Structure-activity Relationship of N-alkyl Gly-boro-Pro Inhibitors of DPP4, FAP, and DPP7." Bioorganic & Medicinal Chemistry Letters 15, 2005, pp. 4239-4242.

Hunziker, D. et al, "Inhibitors of DPP IV-recent advances and structural views", Current Topics in Medicinal Chemistry, 2005, vol. 5 issue 16, pp. 1623-1637.

Huttner, S. et al., "Safety, Tolerability, Pharmacokinetics, and Pharmacodynamics of Single Oral Doses of BI 1356, an Inhibitor of Dipeptidyl Peptidase 4, in Healthy Male Volunteers." Journal of Clinical Pharmacology, 2008, vol. 48, No. 10, pp. 1171-1178.

International Search Report for PCT/EP2007/054204 mailed Mar. 8, 2007.

Inukai, T., "Treatment of Diabetes in Patients for Whom Metformin Treatment is Not Appropriate." Modern Physician, 2008, vol. 28, No. 2, pp. 163-165.

Kharkevich, D. A., "Educational Literature" Pharmacology (1987) Third Edition, Meditsina Press, Moscow pp. 47-48.

Kibbe, A., Editor. Handbook of Pharmaceutical Excipients, Third Edition, Copovidon—pp. 196-197, Date of Revision: Dec. 16, 2008. Mannitol—pp. 424-425, Date of Revision: Feb. 19, 2009, Published in 2009.

Kidney Disease (Nephropathy), Retrieved online May 13, 2013. www.diabetes.org/living-with-diabetes/complications/kidney-disease-nephropathy.html http://www.diabetes.org/living-with-diabetes/complications/kidney-disease-nephropathy.html>

Kim, Kwang-Rok et al., "KR-62436, 6-{2-{2-(5-cyano4,5-dihydropyrazol-1-yl)-2-oxoethylamino}ethylamino}nicotinonitrile, is a novel dipeptidyl peptidase-IV (DDP-IV inhibitor with antihyperglycemic activity" European Journal of Pharmacology 518, 2005, p. 63-70.

Komori, Kiyoshi., "Treatment of Diabetes in Patients for Whom Metforming Treatment is Not Appropriate" Modern Physician (2008) vol. 28, No. 2 pp. 163-165.

Lakatos, P. L. et al., "Elevated Serum Dipeptidyl IV (CD26, EC 3.4.14.5) Activity in Experimental Liver Cirrhosis." European Journal of Clinical Investigation, 2000, vol. 30, No. 9, pp. 793-797. Matsumiya, Teruhiko, et al., "Therapeutic Drugs for Clinicians"

Diagnosis and Treatment (2008) vol. 96, No. 2 pp. 389-390. Mayo Clinic Staff: "Nonalchoholic fatty liver disease: Prevention"

http://www.mayoclinic.com/health/nonalcoholic-fatty-liver-disease/DS00577DSECTION=prevention>.

Nathan, D. et al., "Management of Hyperglycemia in Type 2 Diabetes: A Consensus Algorithm for the Initiation and Adjustment of Therapy." Diabetes Care, Aug. 2006, vol. 29, No. 8, pp. 1963-1972.

OTHER PUBLICATIONS

Nauck, M. A. et al., "Efficacy and Safety of Adding the Dipeptidyl Peptidase-4 Inhibitor Alogliptin to Metformin Therapy in Patients with Type 2 Diabetes Inadequately Controlled with Metformin Monotherapy: A Multicentre, Randomised, Double-Blind, Placebo-Cotrolled Study." Clinical Practice, 2008, vol. 63, No. 1, pp. 46-55. Nauck, M. A. et al., "Efficacy and Safety of the Dipeptidyl Peptidase-4 Inhibitor, Sitagliptin, Compared with the Sulfonylurea, Glipizide, in Patients with Type 2 Diabetes Inaduately Controlled on Metformin alone: A Randomized, Double-Blind, Non-Inferiority Trial." Diabetes Obesity and Metabolism, 2007, vol. 9, No. 2, pp. 194-205

Nielsen, L., "Incretin mimetics and DPP-IV inhibitors for the treatment of type 2 diabetes." Drug Discovery Today, 2005, vol. 10, No. 10, pp. 703-710.

Nihon Ijinpo, Japan Medicinal Journal, 2001, No. 4032, p. 137. Office Action for U.S. Appl. No. 10/695,597 mailed May 2, 2008. Plummer, C.J.G. et al., "The Effect of Melting Point Distributions on DSC Melting Peaks." Polymer Bulletin, 1996, vol. 36, pp. 355-360.

Pratley, R. et al., "Inhibition of DPP-4: a new therapeutic approach for the treatment of type 2 diabetes." Current Medical Research and Opinion, 2007, vol. 23, No. 4, pp. 919-931.

Rosenstock, et al., Sitagliptin Study 019 Groups, Efficacy and safety of the dipeptidyl peptidase-4 inhibitor sitagliptin, Clinical Therapeutics, 2006, vol. 28, Issue 10, p. 1556-1568.

Russell-Jones, D. et al., "Liraglutide vs insulin glargine and placebo in combination with metformin and sulfonylurea therapy in type 2 diabetes mellitus (LEAD-5 met+SU): a randomised controlled trial." Diabetologia, 2009, vol. 52, pp. 2046-2055.

Sarafidis, P. et al., "Cardiometabolic Syndrome and Chronic Kidney Disease: What is the link?" JCMS 2006, 1: p. 58-65.

Schmidt, D. et al., "Fibromatosis of Infancy and Childhood Histology, Ultrastructure and Clinicopathologic Correlation." Zeitschrift für Kinderchirurgie, 1985, vol. 40, No. 1, pp. 40-46.

Shanks, N. et al., Are animal models predictive for humans?, PEHM, Philosophy, Ethics, and Humanaities in Medicine, 4(2), 2009. 1-20.

Shintani, Maki, et al., "Insulin Resistance and Genes" Circulatory Sciences (1997) vol. 17, No. 12 pp. 1186-1188.

Singhal, D. et al., "Drug polymorphism and dosage form design: a practical perspective." Advanced Drug Delivery Reviews, 2004, vol. 56, pp. 335-347.

Targher, G. et al., "Prevalence of Nonalcoholic Fatty Liver Disease and Its Association With Cardiovascular Disease Among Type 2 Diabetic Patients." Diabetes Care, 2007, vol. 30, No. 5, pp. 1212-1218.

Third Party Observation for application No. EP20070728655, May 13, 2013.

Tounyoubyou, "Symposium-19: Future Perspectives on Incretion Therapy in Diabetes." 2008, vol. 51, Suppl. 1, p. S-71, S19-2.

Tradjenta, Highlights of Prescribing Information (revised Sep. 2012).

Tribulova, N. et al. "Chronic Disturbances in NO Production Results in Histochemical and Subcellular Alterations of the Rat Heart." Physiol. Res., 2000, vol. 49, No. 1, pp. 77-88.

Tsujihata, et al., "TAK-875, an orally available G protein-Coupled receptor 40/Free fatty acid receptor 1 Agonist, Enhances Glucose Dependent Insulin Secretion and improves both Postprandial and Fasting hyperglycemic in type 2 Diabetic rats", J. Pharm Exp. 2011, vol. 339, No. 1, p. 228-237.

Uhlig-Laske, B. et al., "Linagliptin, a Potent and Selective DPP-4 Inhibitior, is Safe and Efficacious in Patients with Inadequately Controlled Type 2 Diabetes Despite Metformin Therapy". 535-P Clinical Therapeutics/New Technology—Pharmacologic Treatment of Diabetes or Its Complications, Posters, vol. 58, Jun. 5, 2009, p. A143.

Abstract in English for JP 2002/348279, Dec. 4, 2002. Abstract in English for JP 2003/286287, Oct. 10, 2003.

Adebowale, K.O. et al., "Modification and properties of African yam bean (*Sphenostylis stenocarpa* Hochst. Ex A. Rich.) Harms starch I: Heat moisture treatments and annealing." Food Hydrocolloids, 2009, vol. 23, No. 7, pp. 1947-1957.

Alter, M. et al., "DPP-4 Inhibition on Top of Angiotensin Receptor Bockade Offers a New Therapeutic Approach for Diabetic Nephropathy." Kidney and Blood Pressue Research, 2012, vol. 36, No. 1, pp. 119-130.

Borloo, M. et al. "Dipeptidyl Peptidase IV: Development, Design, Synthesis and Biological Evaluation of Inhibitors." 1994, Universitaire Instelling Antwerpen, vol. 56, pp. 57-88.

Boulton, D.W. et al., "Safety, Tolerability, Pharmacokinetics and Pharmacodynamics of Once-Daily Oral Doses of Saxagliptin for 2 Weeks in Type 2 Diabetic and Healthy Subjects." Diabetes, 2007, Supplement 1, vol. 56, pp. A161.

Chan, J.C. et al., "Safety and efficacy of sitagliptin in patients with type 2 diabetes and chronic renal insufficiency." 2008, Diabetes, Obesity and Metabolism, vol. 10, pp. 545-555.

Chaykovska, L. et al., "Effects of DPP-4 Inhibitors on the Heart in a Rat Model of Uremic Cardiomyopathy." www.plosone.org, 2011, vol. 6, No. 11, p. e27861.

ChemGaroo, "Leaving Group." 1999, Retrieved online: http://www.chemgapedia.de/vsengine/vlu/vsc/en/ch/12/oc/vluorganik/substitution/sn_2/sn 2. vlu/Page/vsc/en/ch/12/oc/substitution/sn_2/abgangsgrupen/abgangsgruppe. vscml.html.

Chisari, A. et al. "Sulphinyl, Sulphonyl, and Sulphonium Groups as Leaving Groups in Aromatic Nucleophilic Substitutions." Journal of the Chemical Society, Perkin Transactions II, 1982, pp. 957-959. Clinical Trials: NCT00309608, "Efficacy and Safety of BI 1356 BS (Linagliptin) in Combination With Metformin in Patients With type2 Diabetes" Boehringer Ingelheim Pharmaceuticals, last updated: Dec. 11, 2013.

Blucotrol XL (glipizide), package insert, Pfizer, Apr. 1, 2002.

Gomez-Perez, et al, "Insulin Therapy:current alternatives", Arch. Med.Res. 36: p. 258-272 (2005).

Graede-Mody, U. et al., "Effect of Renal Impairment on the Pharmacokinetics of the Dipeptidyl Peptidase-4 Inhibitor Linagliptin." Diabetes, Obesity and Metabolism, 2011, pp. 939-946. Groop, P.-H. et al., "Effects of the DPP-4 Inhibitor Linagliptin on Albuminuria in Patients with Type 2 Diabetes and Diabetic Nephropathy." 48th EASD Annual Meeting, Berlin, Abstract 36, Oct. 2012, URI:http://www.abstractsonline.com.

Haluzik, M. et al., "Renal Effects of DPP-4 Inhibitors: A Focus on Microalbuminuria." International Journal of Endocrinology, 2013, vol. 35, No. 6, pp. 1-7.

Hansen, H. et al., "Co-Administration of the DPP-4 Inhibitor Linagliptin and Native GLP-1 Induce Body Weight Loss and Appetite Suppression." 73rd Annual Meeting Science Session, ADA, Chicago, Jun. 21, 2013.

Headland, K. et al., "The Effect of Combination Linagliptin and Voglibose on Glucose Control and Body Weight." 73rd Annual Meeting Science Session, ADA, Chicago, Jun. 21, 2013.

Heihachiro, A. et al., "Synthesis of Prolyll Endopeptidase Inhibitors and Evaluation of Their Structure-Activity Relationships: In Vitro Inhibition of Prolyl Endopeptidase from Canine Brain." 1993, Chemical and Pharmaceutical Bulletin, vol. 41, pp. 1583-1588.

Hocher, B. et al., "Renal and Caridae Effects of DPP-4 Inhibitors—from Preclinical Development to Clinical Research." Kidney & Blood Pressue Research, 2012, vol. 36, No. 1, pp. 65-84.

Holman, et al., "Additon of Biphasic, prandial, or basal insulin to oral therapy in type 2 diabetes", N. England Journal Medicine, p. 1716-1730, 2007.

International Search Report and Written Opinion for PCT/EP2011/057256 dated Jul. 22, 2011.

International Search Report and Written Opinion for PCT/EP2012/077024 mailed Feb. 19, 2013.

International Search Report and Written Opinion for PCT/EP2013/059828 mailed Aug. 6, 2013.

International Search Report and Written Opinion for PCT/EP2013/059831 mailed on Aug. 9, 2013.

International Search Report and Written Opinion for PCT/EP2013/060311 mailed Aug. 9, 2013.

OTHER PUBLICATIONS

International Search Report and Written Opinion for PCT/EP2013/060312 mailed on Sep. 4, 2013.

International Search Report and Written Opinion for PCT/EP2013/070978 mailed on Oct. 31, 2013.

International Search Report for PCT/EP2013/060309 mailed Aug. 9, 2013.

Johansen, O. E. et al., "Cardiovascular with linagliptin in patients with type 2 diabetes mellitus: a pre-specified, prospective, and adjudicated meta-analysis of a phase 3 programme." Cardiovascular Diabetology, Biomed Central, 2012, vol. 11, No. 1, pp. 1-10.

Johansen, O.E. et al., "b-cell Function in Latnet Autoimmune Diabetes in Adults (LADA) Treated with Linagliptin Versus Glimepiride: Exploratory Results from a Two Year Double-Blind, Randomized, Controlled Study." www.abstractsonline.com, Jun. 10, 2012. XP-002708003.

John Hopkins Children's Center, "Liver Disorders and Diseases." Retrieved online May 26, 2014 http://www.hopkinschildrens.org/non-alcoholic-fatty-liver-disease.aspx>.

Kroller-Schön, S. et al., "Glucose-independent Improvement of Vascular Dysfunction in Experimental Sepsis by Dipeptidyl Peptidase-4 Inhibition." Cardiovascular Research, 2012, vol. 96, No. 1, pp. 140-149.

Lee Jones, K. et al., "Effect of Metformin in Pediatric Patients With Type 2 Diabetes." Diabetes Care, 2002, vol. 25, No. 1, pp. 89-94. Lovshin, J.A. et al., "Incretin-based therapies for type 2 diabetes mellitus." Nature Reviews Endocrinology, 2009, vol. 5, pp. 262-269

Nielsen, L., "Incretin Mimetics and DPP-IV Inhibitors for the Treatment of Type 2 Diabetes." DDT, 2005, vol. 10, No. 10, pp. 703-710

Rosenbloom, et al., "Type 2 Diabetes mellitus in the child and adolescent", Pediatric Diabetes, 2008, p. 512-526.

Sharkovska, Y., et al., "DPP-4 Inhibition with Linagliptin Delays the Progression of Diabetic Nephropathy in db/db Mice." 48th EASD Annual Meeting, Berlin, Abstract 35, Oct. 2012, URL:http://www.abstractsonline.com.

Silverman, G. et al., "Handbook of Grignard Reagents." 1996, Retrieved online: http://books.google.com/books?id=82CaxfY-uNkC&printsec=frontcover&

dq=intitle:Handbook+intitle:of+intitle:Grignard+intitle:Reagents&hl=en&sa=X&ei=g06GU5SdOKngsATphYCg&

ved=0CDYQ6AEwAA#v=onepgae&q&f=false>.

Sortino, M.A. et al., "Linagliptin: a through characterization beyond its clinical efficacy." Frontiers in Endocrinology, 2013, vol. 4, Article 16, pp. 1-9.

Thornber, C.W., "Isosterism and Molecular Modification in Drug Design." Chemical Society Reviews, 1979, pp. 563-580.

Tsuprykov, O. et al., Linagliptin is as Efficacious as Telmisartan in Preventing Renal Disease Progression in Rats with 5/6 Nephrectomy, 73rd Annual Meeting Science Session, ADA, Chicago, Jun. 2013 URI:http://www.abstractonline.com/.

Wertheimer, et al., "Drug Delivery Systems improve pharmaceutical profile and faciliate medication adherence", Adv. Therapy 22: p. 559-577 (2005).

Youssef, S. et al., "Purines XIV. Reactivity of 8-Promo-3,9-dimethylxanthine Towards Some Nucleophilic Reagents." Journal of Heterocyclic Chemistry, 1998, vol. 35, pp. 949-954.

Al-Masri, I.M. et al., "Inhibition of dipeptidyl peptidase IV (DPP IV) is one of the mechanisms explaining the hypoglycemic effect of berberine." Journal of Enzyme Inhibition and Medicinal Chemistry, 2009, vol. 24, No. 5, pp. 1061-1066.

Al-Masri, Journal of Enzyme Inhibition and Medicinal Chemistry, "Inhibition of dipeptyl peptidase IV (DPP iv) is one of the mechanisms explaining hypoglycemic effect of berberine", 2009.

Forst, T. et al., "The oral DPP-4 inhibitor linagliptin significantly lowers HbA1c after 4 weeks of treatment in patients with type 2 diabetes mellitus." Diabetes, Obesity and Metabolism, 2011, vol. 13, pp. 542-550.

Hashida, Mitsuru, "Strategies for designing and developing oral administration formulations." Yakuji-Jiho, Inc., 1995, pp. 50-51. Leibovitz, Cardiovascular Diabetology, Sitagliptin Treatment in diabetes patients presenting with acute coronary syndrome: results from the Acute Coronary Syndrome Israeli Survey, 2013.

Leibovitz, E. et al., "Sitagliptin pretreatment in diabetes patients presenting with acute coronary syndrome: results from the Acute Coronary Syndrome Israeli Survey (ACSIS)." Cardiovascular Diabetology, 2013, vol. 12, No. 1, pp. 1-7.

Lim, S. et al., "Effect of a Dipeptidyl Peptidase-IV Inhibitor, Des-Fluoro-Sitagliptin, on Neointimal Formation after Balloon Injury in Rats." Plos One, 2012, vol. 7, No. 4, pp. 1-11.

Lim, Seoul National Univ. Bundang Hospital, Effect of a Dipeptyl Peptidase-IV Inhibitor, Des-Fluoro Sitagliptin, on Neointimal Formation after Balloon Injury in Rats, 2012, vol. 7, Issue 4, Plos One. Lyssenko, V. et al., "Mechanisms by which common variants in the TCF7L2 gene increase risk of type 2 diabetes." The Journal of Clinical Investigation, 2007, vol. 117, No. 8, pp. 2155-2163.

Matsumiya, Tokyo Medical University, Department of Pharmacology, Diagnosis and Therapy, vol. 96, No. 2, 2008, pp. 389-390. McNay, David E.G. et al., "High fat diet causes rebound weight gain." Molecular Metabolism, 2013, vol. 2, pp. 103-108.

Iwamoto Insulin Glargine, Nippon Rinsho 60(9):503-515.

Partial International Search Report for PCT/EP2014/060160, mailing date Jun. 30, 2014.

Pearson, E. R. et al., "Variation in TCF7L2 Influences Therapeutic Response to Sulfonylureas." Diabetes, 2007, vol. 56, pp. 2178-2182.

Prescribing Information, Package insert for Leprinton tablets 100mg, Manufacturer: Tatsumi Kagaku Co., Ltd., Mar. 2003.

Radermecker, Regis et al., "Lipodystrophy Reactions to Insulin." American Journal of Clinical Dermatology, 2007, vol. 8, pp. 21-28. Rask-Madsen, C. et al., "Podocytes lose their footing." Nature, 2010, vol. 468, pp. 42-44.

Sheperd, Todd M. et al., "Efective management of obesity." The Journal of Family Practice, 2003, vol. 52, No. 1, pp. 34-42.

Shu, L. et al., "Decreased TCF7L2 protein levels in type 2 diabetes mellitus correlate with downregulation of GIP- and GLP-1 receptors and impaired beta-cell function." Human Molecular Genetics, 2009, vol. 18. No. 13, pp. 2388-2399.

vol. 18, No. 13, pp. 2388-2399. Shu, L. et al., "Transcription Factor 7-Like 2 Regulates B-Cell Survival and Function in Human Pancreatic Islets." Diabetes, 2008, vol. 57, pp. 645-653.

Suzuki, Y. et al., "Carbon-Carbon Bond Cleavage of a-Hydroxybenzylheteroarenes Catalyzed by Cyanide Ion: Retro-Benzoin Condensation Affords Ketones and Heteroarenes and Benzyl Migration Affords Benzylheteroarenes and Arenecarbaldehydes." Chemical Pharmaceutical Bulletin, 1998, vol. 46(2), pp. 199-206.

Zimdahl, H. et al., "Influence of TCF7L2 gene variants on the therapeutic response to the dipeptidylpeptidase-4 inhibitor linagliptin." Diabetologia, 2014, vol. 57, pp. 1869-1875.

Ahren, Bo "Novel combination treatment of type 2 diabetes DPP-4 inhibition + metformin." Vascular Health and Risk Management, 2008, vol. 4, No. 2, pp. 383-394.

Baetta, R. et al., "Pharmacology of Dipeptidyl Peptidase-4 Inhibitors." Drugs, 2011, vol. 71, No. 11, pp. 1441-1467.

Blech, S. et al., "The Metabolism and Disposition of the Oral Dipeptidyl Peptidase-4 Inhibitor, Linagliptin, in Humans", Drug Metabolism and Disposition, 2010, vol. 38, No. 4, p. 667-678.

Cheon, et al., Biochemical Pharmacology, "Inhibition of dipeptidyl IV by novel inhibitors with pyrazolidine scaffold", 2005, vol. 70, p. 22-29.

Clinical Trials, No. NCT00309608, "Efficacy and Safety of BI 1356 BS in Combination with Metformin in Patients With type2 Diabetes" 2009, pp. 1-3.

Craddy, P. et al., "Comparative Effectiveness of Dipeptidylpeptidase-4 Inhibitors in Type 2 Diabetes: A Systematic Review and Mixed Treatment Comparison." Diabetes Therapy, 2014, vol. 5, No. 1, pp. 1-41.

Crowe, E. et al., "Early identification and management of chronic kidney disease: summary of NICE guidance." British Medical Journal, 2008, vol. 337, pp. 812-815.

OTHER PUBLICATIONS

Drucker, Daniel J., "Dipeptidyl Peptidase-4 Inhibition and the Treatment of Type 2 Diabetes." Diabetes Care, 2007, vol. 30, No. 6, pp. 1335-1343.

Gallwitz, B. et al., "2-year efficacy and safety of linagliptin compared with glimepiride in patients with type 2 diabetes inadequately controlled on metformin: a randomised, double-blind, non-inferiority trial." Lancet, 2012, vol. 380, pp. 475-483.

Greischel, et al., Drug Metabolism and Deposition, "The Dipeptidyl Peptidase-4 Inhibitor Linagliptin Exhibits Time-and DPSE-Dependent Localization in Kidney, Liver, and Intestine after Intravenous Dosing: Results from High Resolution Autoradiography in Rats", 2010, vol. 38, No. 9, p. 1443-1448.

Guglielmi, C. et al., "Latent autoimmune diabetes in the adults (LADA) in Asia: from pathogenesis and epidemiology to therapy." Diabetes/Metabolism Research and Reviews, 2012, vol. 28, Supplement 2, pp. 40-46.

Gupta, V. et al., "Choosing a Gliptin." Indian Journal of Endocrinology and Metabolism, 2011, vol. 15, No. 4, pp. 298-308.

Heise, et al., Diabetes, Obesity and Metabolism, "Pharmacokinetics, pharmacokinetics and tolerability of mutilple oral doses of linagliptin, a dipeptidyl peptidase-4 inhibitor in male type 2 diabetes patients", 2009, vol. 11, No. 8, p. 786-794.

Hocher, B. et al., "The novel DPP-4 inhibitors linagliptin and BI 14361 reduce infarct size after myocardial ischemia/reperfusion in rats." International Journal of Cardiology, 2013, vol. 167, pp. 87-93. Hull, R. et al., "Nephrotic syndrome in adults." British Medical Journal, 2008, vol. 336, pp. 1185-1190.

Isomaa, B. et al., "Cardiovascular Morbidity and Mortality Associated With the Metabolic Syndrome." Diabetes Care, 2001, vol. 24, No. 4, pp. 683-689.

Kendall, D. M. et al., "Incretin Mimetics and Dipeptidyl Peptidase-IV Inhibitors: A Review of Emerging Therapies for Type 2 Diabetes." Diabetes Technology & Therapeutics, 2006, vol. 8, No. 3, pp. 385-398.

Klein, T. et al., "Linagliptin alleviates hepatic steatosis and inflammation in a mouse model of non-alcoholic steatohepatitis." Medical Molecular Morphology, 2014, vol. 47, pp. 137-149.

Konstantinou, D. M. et al., "Pathophysiology-based novel pharmacotherapy for heart failure with preserved ejection fraction." Pharmacology & Therapeutics, 2013, vol. 140, No. 2, pp. 156-166. Lakatos P. L. et al., "Elevated serum dipeptidyl peptidase IV (CD26, EC 3.4.14.5) activity in patients with primary biliary cirrhosis." Journal of Hepatol, 1999, vol. 30, p. 740.

Linagliptin Monograph, Published by VACO PBM-SHG US Veteran's Administration, 2011, pp. 1-17.

Naik, R. et al., "Latent Autoimmune Diabetes in Adults." The Journal of Clinical Endocrinology and Metabolism, 2009, vol. 94, No. 12, pp. 4635-4644.

National Program for Care Guidelines, "Type 2 Diabetes mellitus." 2002, First Edition, pp. 1-50.

Poudel, Resham R., "Latent autoimmune diabetes of adults: From oral hypoglycemic agents to early insulin." Indian Journal of Endocrinology and Metabolism, 2012, vol. 16, Supplement 1, pp. S41-S46.

Schillinger, M. et al., "Restenosis after percutaneous angioplasty: the role of vascular inflammation." Vascular Health and Risk Management, 2005, vol. 1, No. 1, pp. 73-78.

Schurmann, C. et al., "The Dipeptidyl Peptidase-4 Inhibitor Linagliptin Attenuates Inflammation and Accelerates Epithelialization in Wounds of Diabetic ob/ob Mice." The Journal of Pharmacology and Experimental Therapeutics, 2012, vol. 342, No. 1, pp. 71-80.

Standl, E. et al., "Diabetes and the Heart." Diabetes Guidelines (DDG), 2002, pp. 1-25.

Tadayyon, M. et al., "Insulin sensitisation in the treatment of Type 2 diabetes." Expert Opinion Investigative Drugs, 2003, vol. 12, No. 3, pp. 307-324.

Takeda Press Release: "Voglibose (BASEN) for the prevention of type 2 diabetes mellitus: A Randomized, Double-blind Trial in

Japanese Subjects with Impaired Glucose Tolerance." 2008, Retrieved online Jul. 6, 2015. https://www.takeda.com/news/2008/20080526_3621.html.

Taskinen, M.-R. et al., "Safety and efficacy of linagliptin as add-on therapy to metformin in patients with type 2 diabetes: a randomized, double-blind, placebo-controlled study." Diabetes, Obesity and Metabolism, 2011, vol. 13, pp. 65-74.

Vichayanrat, A. et al., "Efficacy and safety of voglibose in comparison with acarbose in type 2 diabetic patients." Diabetes Research and Clinical Practice, 2002, vol. 55, pp. 99-103.

Vickers, 71st Scientific Session of the American Diabetes Association, "The DPP-4 inhibitor linagliptin is weight neutral in the DIO rat but inhibits the weight gain of DIO animals withdrawn from exenatide", vol. 60, Jul. 2011.

Weber, Ann E., "Dipeptidyl Peptidase IV Inhibitors for the Treatment of Diabetes." Journal of Medicinal Chemistry, 2004, vol. 47, pp. 4135-4141.

WebMD, Autoimmune Diseases: What Are They? Who Gets Them? "What Are Autoimmune Disorders?" 2015, pp. 1-3. Retrieved online Jul. 9, 2015. http://www.webmd.com/a-to-z-guides/autoimmune-diseases.

Witteles, R. M. et al., "Dipeptidyl Peptidase 4 Inhibition Increases Myocardial Glucose Uptake in Nonischemic Cardiomyopathy." Journal of Cardiac Failure, 2012, vol. 18, No. 10, pp. 804-809.

Yamagishi, S. et al., "Pleiotropic Effects of Glucagon-like Peptide-1 (GLP-1)-Based Therapies on Vascular Complications in Diabetes." Current Pharmaceutical Design, 2012, vol. 17, pp. 4379-4385.

Zhimei, Xiao et al., "Study progression of oral drugs for treatment of type II diabetes." Drug Evaluation, 2004, vol. 1, No. 2, pp. 138-143.

Anonymous, Clinicaltrials.gov, 2006, No. NCT00309608, "Efficacy and Safety of BI 1356 BS in Combination with Metformin in Patients With type2 Diabetes" pp. 1-3.

Greischel, et al., Drug Metabolism and Deposition, "The Dipeptidyl Peptidase-4 Inhibitor Linagliptin Exhibits Timeand DPSE-Dependent Localization in Kidney, Liver, and Intestine after Intravenous Dosing: Results from High Resolution Autoradiography in Rats", 2010, vol. 38, No. 9, p. 1443-1448.

Lakatos, P. L. et al., "Elevated serum dipeptidyl peptidase IV (CD26, EC 3.4.14.5) activity in patients with primary biliary cirrhosis." Journal of Hepatol, 1999, vol. 30, p. 740.

Witteles, R. M. et al., "Pipeptidyl Peptidase 4 Inhibition Increases Myocardial Glucose Uptake in Nonischemic Cardiomyopathy." Journal of Cardiac Failure, 2012, vol. 18, No. 10, pp. 804-809. Abstract for AU 2003280680, Jun. 18, 2004.

Abstract for AU 2009224546, Sep. 17, 2009.

Abstract in English for DE19705233, Aug. 13, 1998.

ACTOS Prescribing Information, 1999, pp. 1-26.

Ahren, B. et al., "Twelve- and 52-Week Efficacy of the Dipeptidyl Peptidase IV Inhibitor LAF237 in Metformin-Treated Patients With Type 2 Diabetes." Diabetes Care, 2004, vol. 27, No. 12, pp. 2874-2880.

American Association of Clinical Endocrinologists, "Medical Guidelines for Clinical Practice for the Management of Diabetes Mellitus." Endocrine Practice, 2007, col. 13, Suppl. 1, pp. 1-68.

Banker, Gilbert S., "Prodrugs." Modem Pharmaceutics Third Edition, Marcel Dekker, Inc., 1996, p. 596.

Beauglehole, Anthony R., "N3-Substituted Xanthines as Irreversible Adenosine Receptor Antagonists." Ph.D. Thesis, Deakin University, Australia, 2000, pp. 1-168.

Canadian Diabetes Association, "Pharmacologic Management of Type 2 Diabetes." Canadian Journal of Diabetes, 2003, vol. 27, Suppl. 2, pp. S37-S42.

Chiasson, J.-L. et al., "The Synergistic Effect of Miglitol Plus Metformin Combination Therapy in the Treatment of Type 2 Diabetes." Diabetes Care, 2001, vol. 24, No. 6, pp. 989-994.

Clinical Trial Protocol, "A Randomised, Double-blind, Placebocontrolled, Five Parallel Groups Study Investigating the Efficacy and Safety of BI 1356 BS." Boehringer Ingelheim Pharmaceuticals, last updated on Jun. 24, 2014.

Clinical Trial, NCT00622284, clinicaltrials.gov, updated Feb. 22, 2008, pp. 1-3.

OTHER PUBLICATIONS

Clinical Trials NCT00601250, clinicaltrials.gov, Jan. 25, 2008, pp. 1-3.

Clinical Trials: NCT00103857, "A Multicenter, Randomized, Double-Blind Factorial Study of the Co-Administration of MK0431 and Metformin in Patients With Type 2 Diabetes Mellitus Who Have Inadequate Glycemic Control" last updated on Apr. 27, 2015, pp. 1-4.

Clinical Trials: NCT00309608, "Efficacy and Safety of BI 1356 BS (Linagliptin) in Combination With Metformin in Patients With type2 Diabetes" Boehringer Ingelheim Pharmaceuticals, last updated on Jun. 24, 2014, pp. 1-3.

Colorcon, "Lactose Replacement with Starch 1500 in a Direct Compression Formula." 2005, pp. 1-4.

Colorcon, "Reducing Coated Tablet Defects from Laboratory through Production Scale: Performance of Hypromellose or Polyvinyl Alcohol-Based Aqueous Film Coating Systems." Opadry II, 2009, pp. 1-7.

Deacon, Carolyn F., "Dipeptidyl peptidase 4 inhibition with sitagliptin: a new therapy for Type 2 diabetes." Expert Opinion on Investigational Drugs, 2007, vol. 16, No. 4, pp. 533-545.

Dittberner, S. et al., "Determination of the absolute bioavailability of BI 1356, a substance with non-linear pharmacokinetics, using a population pharmacokinetic modeling approach." Abstracts of the Annual Meeting of the Population Approach Group in Europe, 2007.

Dugi, K. et al., "Safety, tolerability, pharmacokinetics, and pharmacodynamics of BI 1356, a novel DPP-IV inhibitor with a wide therapeutic window." Diabetic Medicine, 2006, vol. 23, Suppl. 4, p. 300, p. 821.

EMEA: European Medicines Agency, "Galvus (vildagliptin)" Retrieved online on Jan. 21, 2016, pp. 1-6.

Feng, J. et al., "Discovery of Alogliptin: A Potent, Selective, Bioavailable, and Efficacious Inhibitor of Dipeptidyl Peptidase IV." Journal of Medicinal Chemistry, 2007, vol. 50, No. 10, pp. 2297-2300.

Flatt, P.R. et al., "Dipeptidyl peptidase IV (DPP IV) and related molecules in type 2 diabetes." Frontiers in Bioscience, 2008, vol. 13, pp. 3648-3660.

Gall, "Prevalence of micro-and macroalbuminuria, arterial hypertension, retinopathy and large vessel disease in European type 2 (non-insulin dependent) diabetic patients", Diabetologia (1991) 655-661.

Gallwitz, B., "Safety and efficacy of linagliptin in type 2 diabetes patients with common renal and cardiovascular risk factors." Therapeutic Advances in Endocrinology and Metabolism, 2013, vol. 4, No. 3, pp. 95-105.

Galvus (Vildagliptin) Scientific Discussion, EMEA, 2007, pp. 1-34. Garber, A.J. et al., "Simultaneous glyburide/metformin therapy is superior to component monotherapy as an initial pharmacological treatment for type 2 diabetes." Diabetes, Obesity and Metabolism, 2002, vol. 4, pp. 201-208.

Glucophage® Prescribing Information, 2001, pp. 1-7.

Goodarzi, M.O. et al., "Metformin revisited: re-evaluation of its properties and role in the pharmacopoeia of modem antidiabetic agents." Diabetes, Obesity and Metabolism, 2005, vol. 7, pp. 654-665.

Gwaltney, S.L. II et al., "Inhibitors of Dipeptidyl Peptidase 4." Annual Reports in Medicinal Chemistry, 2005, vol. 40, pp. 149-165. Halimi, "Combination treatment in the management of type 2 diabetes: focus on vildagliptin and metformin as a single tablet", Vascular Health and Risk Management, 2008 481-92.

He, Y.L. et al., "The Influence of Renal Impairment on the Pharmacokinetics of Vildagliptin." Clinical Pharmacology & Therapeutics, 2007, vol. 81, Suppl. 1, Abstract No. PIII-86.

Rinke, S.A. et al., "Metformin Effects on Dipeptidylpeptidase IV Degradation of Glucagon-like Peptide-1." Biochemical and Biophysical Research Communications, 2002, vol. 291, No. 5, pp. 1302-1308.

Hinke, S.A. et al., "On Combination Therapy of Diabetes With Metformin and Dipeptidyl Peptidase IV Inhibitors." Diabetes Care, 2002, vol. 25, No. 8, pp. 1490-1492.

Hinnen, D. et al., "Incretin Mimetics and DPP-IV Inhibitors: New Paradigms for the Treatment of Type 2 Diabetes." Journal of the American Board of Family Medicine, 2006, vol. 19, No. 6, pp. 612-620.

International Search Report for PCT/EP2014/060160 mailed Nov. 8, 2014, pp. 1-5.

Inzucchi, Silvio E., "Oral Antihyperglycemic Therapy for Type 2 Diabetes." The Journal of the American Medical Association, 2002, vol. 287, No. 3, pp. 360-372.

Janumet Prescribing Information, revised Jan. 2008, pp. 1-22.

Januvia Prescribing Information and Product Label, 2006, pp. 1-36. Kiraly, K. et al., "The dipeptidyl peptidase IV (CD26, EC 3.4.14.5) inhibitor vildagliptin is a potent antihyperalgesic in rats by promoting endomorphin-2 generation in the spinal cord." European Journal of Pharmacology, 2011, vol. 650, pp. 195-199.

Kirpichnikov, D. et al., "Metformin: An Update" Annals of Internal Medicine, 2002, vol. 137, No. 1, pp. 25-33.

Knowler, W.C. et al., "Reduction in the Incidence of Type 2 Diabetes with Lifestyle Intervention or Metformin." The New England Journal of Medicine, 2002, vol. 346, No. 6, pp. 393-403. Lachman, L. et al., "The Theory and Practice of Industrial Pharmacy." Varghese Publishing House, Third Edition, 1987, pp. 190-194.

Lindsay, J.R. et al., "Inhibition of dipeptidyl peptidase IV activity by oral metformin in Type 2 diabetes." Diabetic Medicine, 2005, vol. 22, pp. 654-657.

Lu, "High prevlaence of albuminuria in population based patients diagnosed with type 2 diabetes in the Shanghai downtown", Diabestes Research and Clinical Practice (2007) 184-192.

Mathieu, C. et al., "Antihyperglycaemic therapy in elderly patients with type 2 diabetes: potential tole of incretin mimetics and DPP-4 inhibitors." International Journal of Clinical Practice, 2007, vol. 61, Suppl. 154, pp. 29-37.

Merck Manual of Diagnosis and Therapy: "Obesity." 1999, 17th Edition, Chapter 5, pp. 58-62.

Mikhail, Nasser, "Incretin mimetics and dipeptidyl peptidase 4 inhibitors in clinical trials for the treatment of type 2 liabetes." Expert Opinion on Investigational Drugs, 2008, vol. 17, No. 6, pp. 845-853.

Novartis AG, Investor Relations Release, "Galvus, a new oral treatment for type 2 diabetes, receives positive opinion recommending European Union approval." Securities and Exchange Commission, Form 6-K, 2007, pp. 1-4.

Pietruck, F. et al., "Rosiglitazone is a safe and effective treatment option of new-onset diabetes mellitus after renal transplantation." Transplant International, 2005, vol. 18, pp. 483-486.

Sulkin, T.V. et al., "Contraindications to Metformin Therapy in Patients With NIDDM." Diabetes Care, 1997, vol. 20, No. 6, pp. 925-928.

Turner, R.C. et al., "Glycemic Control With Diet, Sulfonylurea, Metformin, or Insulin in Patients With Type 2 Diabetes Mellitus Progressive Requirement for Multiple Therapies (UKPDS 49)" The Journal of the American Medical Association, 1999, vol. 281, No. 21, pp. 2005-2012.

Van Heek, M. et al., "Ezetimibe, a Potent Cholesterol Absorption Inhibitor, Normalizes Combined Dyslipidemia in obese Hyperinsulinemic Hamsters." Diabetes, 2001, vol. 50, pp. 1330-1335

Vincent, S.H. et al., "Metabolism and Excretion of the Dipeptidyl Peptidase 4 Inhibitor [14C]Sitagliptin in Humans." Drug Metabolism and Disposition, 2007, vol. 35, No. 4, pp. 533-538.

Yale, Jean-Francois, "Oral Antihyperglycemic Agents and Renal Disease: New Agents, New Concepts." Journal of the American Society of Nephrology, 2005, vol. 16, Suppl. 1, pp. S7-S10.

Yap, W.S. et al., "Review of management of type 2 diabetes mellitus." Journal of Clinical Pharmacy and Therapeutics, 1998, vol. 23, pp. 457-465.

OTHER PUBLICATIONS

Yokoyama< "Prevalence of albumineria and renal insufficiency and associated clinical factors in type 2 diabetes: The Japan Diabetes clinical data Management study(JDDM15)" Nephrol Dial Transplant (2009) 24: 1212-1219 Advance Access Pub 2008.

Zander, M. et al., "Additive Glucose-Lowering Effects of Glucagon-Like Peptide-1 and Metformin in Type 2 Diabetes." Diabetes Care, 2001, vol. 24, No. 4, pp. 720-725.
Zerilli, T. et al., "Sitagliptin Phosphate: A DPP-4 Inhibitor for the

Zerilli, T. et al., "Sitagliptin Phosphate: A DPP-4 Inhibitor for the Treatment of Type 2 Diabetes Mellitus." Clinical Therapeutics, 2007, vol. 29, No. 12, pp. 2614-2634.

* cited by examiner

Baseline HbA1c values for the whole patient population of the studies (full analysis set, FAS), for the subpopulation for which genetic analyses are performed (full analysis set for pharmacogenetic analyses, FASG), as well as for the subgroups defined by SNP rs7903146 in TCF7L2 genotypes (CC, CT, TT) of this subpopulation.

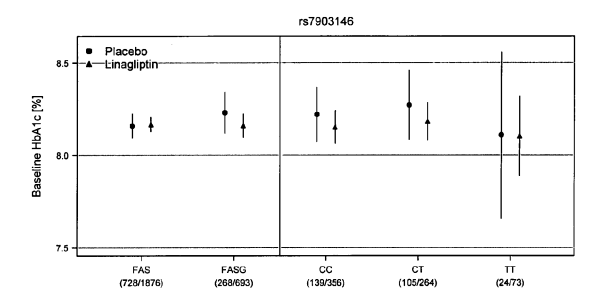
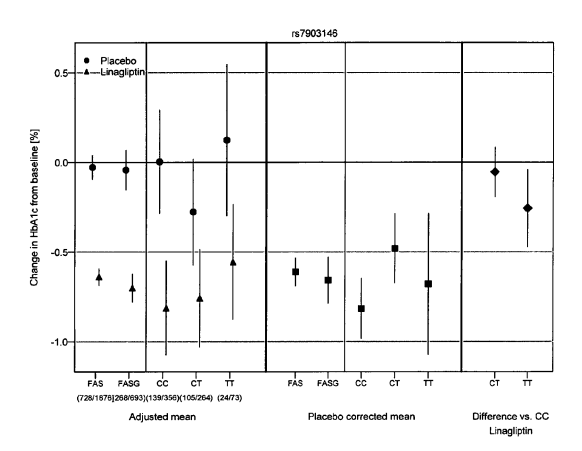


Figure 2 Association of SNP rs7903146 in TCF7L2 with linagliptin response



TREATMENT OF GENOTYPED DIABETIC PATIENTS WITH DPP-IV INHIBITORS SUCH AS LINAGLIPTIN

TECHNICAL FIELD OF THE INVENTION

The invention describes DPP-4 inhibitors, pharmaceutical compositions or combinations comprising a DPP-4 inhibitor as defined herein and optionally one or more other active substances, for use in methods of treatment or prevention as described herein, such as e.g. of one or more conditions selected from type 1 diabetes mellitus, type 2 diabetes mellitus, impaired glucose tolerance, impaired fasting blood glucose and hyperglycemia inter alia. In a particular embodiment, the therapeutic and/or preventive methods of this invention comprise the step of identifying a patient being susceptible to the treatment and/or prevention, said identifying comprising testing whether the patient has variation(s) in one or more genes associated with metabolic diseases 20 (e.g. whether the patient is of a TCF7L2 risk genotype as described herein) or whether the patient is of respective wild-type genotype (e.g. whether the patient is of TCF7L2 wild genotype as described herein), and the further step of administering such DPP-4 inhibitor, pharmaceutical compo- 25 sition or combination to the patient determined as being susceptible.

Further, in one embodiment, the usability of a DPP-4 inhibitor, a pharmaceutical composition, combination or medicament each as described herein for a therapeutic 30 and/or preventive method or use according this invention in a patient who has variation(s) in one or more genes associated with metabolic diseases (such as e.g. a TCF7L2 risk genotype patient as described herein) is contemplated.

TCF7L2 risk genotype patients according to this invention include, without being limited, patients (particularly type 2 diabetes patients) harboring genetic risk variants in the gene TCF7L2 and suffering often from the pathological influences thereof, particularly associated with the risk T-allele of TCF7L2 rs7903146, such as patients harboring the 40 TCF7L2 rs7903146 CT heterozygous risk genotype or patients harboring the TCF7L2 rs7903146 TT homozygous high risk genotype.

Further, in another embodiment, the usability of a DPP-4 inhibitor, a pharmaceutical composition, combination or 45 medicament each as described herein for a therapeutic and/or preventive method or use according this invention in a patient who carries the TCF7L2 wild genotype, particularly the TCF7L2 rs7903146 CC wild genotype, is contemplated.

Moreover, the present invention provides a diagnostic method for identifying a subject (particularly a type 2 diabetes patient) statistically more likely to have a favorable response (e.g. in achieving glycemic control, such as change in HbA1c) to the administration of a therapeutically effec- 55 tive amount of a DPP-4 inhibitor, optionally in combination with one or more other active substances (e.g. antidiabetics), said method comprising determining whether the subject is either of TCF7L2 risk genotype (particularly TCF7L2 rs7903146 CT or TT risk genotype) or of TCF7L2 wild 60 genotype (particularly TCF7L2 rs7903146 CC wild genotype), wherein the subject being of TCF7L2 rs7903146 CC homozygous wild genotype (and, to a lesser extent, the subject being of TCF7L2 rs7903146 CT heterozygous risk genotype) has an increased likelihood of favorable response to the administered DPP-4 inhibitor relative to a subject of TCF7L2 rs7903146 TT homozygous risk genotype.

2

Furthermore the invention describes a method for preventing, slowing progression of, delaying, or treating a metabolic disorder;

for improving glycemic control and/or for reducing of fasting plasma glucose, of postprandial plasma glucose and/or of glycosylated hemoglobin HbA1c;

for preventing, slowing, delaying or reversing progression from impaired glucose tolerance, impaired fasting blood glucose, insulin resistance and/or from metabolic syndrome to type 2 diabetes mellitus;

for preventing, slowing progression of, delaying or treating of a condition or disorder selected from the group consisting of complications of diabetes mellitus;

for reducing body weight and/or body fat or preventing an increase in body weight and/or body fat or facilitating a reduction in body weight and/or body fat;

for preventing or treating the degeneration of pancreatic beta cells and/or for improving and/or restoring or protecting the functionality of pancreatic beta cells and/or restoring the functionality of pancreatic insulin secretion:

for preventing, slowing, delaying or treating diseases or conditions attributed to an abnormal accumulation of liver or ectopic fat; or

for maintaining and/or improving the insulin sensitivity and/or for treating or preventing hyperinsulinemia and/ or insulin resistance;

for preventing, slowing progression of, delaying, or treating new onset diabetes after transplantation (NODAT) and/or post-transplant metabolic syndrome (PTMS);

for preventing, delaying, or reducing NODAT and/or PTMS associated complications including micro- and macrovascular diseases and events, graft rejection, infection, and death;

for treating hyperuricemia and hyperuricemia associated conditions:

in patients in need thereof, for example in those patients (particularly type 2 diabetes mellitus patients) who have variation(s) in one or more genes associated with metabolic diseases (such as e.g. in a TCF7L2 risk genotype patient as described herein) or in those patients which are of respective wild-type genotype (such as e.g. in a TCF7L2 wild genotype as described herein), wherein said method comprises

testing the patient whether he/she has variation(s) in one or more genes associated with metabolic diseases (e.g. whether he/she is of a TCF7L2 risk genotype as described herein) or whether the patient is of respective wild-type genotype (e.g. whether the patient is of TCF7L2 wild genotype as described herein), and

administering a DPP-4 inhibitor as defined hereinafter (preferably linagliptin), optionally in combination with one or more other active substances.

In addition, the present invention describes the use of a DPP-4 inhibitor for the manufacture of a medicament for use in a method as described hereinbefore and hereinafter.

In addition, the present invention describes a DPP-4 inhibitor for use in a therapy of a patient (particularly human type 2 diabetes patient) as described hereinbefore and hereinafter

In addition, the present invention describes a DPP-4 inhibitor for use in a treatment or prevention of a (particularly metabolic) disease, disorder or condition (particularly diabetes, especially type 2 diabetes, and conditions related thereto, such as e.g. diabetic complications) as described hereinbefore and hereinafter.

The invention also describes a use of a pharmaceutical composition or combination according to this invention for

the manufacture of a medicament for use in a method as described hereinbefore and hereinafter.

The invention also relates to the DPP-4 inhibitors as defined herein for use in a method as described hereinbefore and hereinafter, said method comprising administering the 5 DPP-4 inhibitor, optionally in combination with one or more other active substances (e.g. which may selected from those mentioned herein), to the patient.

BACKGROUND OF THE INVENTION

Type 2 diabetes is an increasingly prevalent disease that due to a high frequency of complications leads to a significant reduction of life expectancy. Because of diabetes-associated microvascular complications, type 2 diabetes is 15 currently the most frequent cause of adult-onset loss of vision, renal failure, and amputations in the industrialized world. In addition, the presence of type 2 diabetes is associated with a two to five fold increase in cardiovascular disease risk.

After long duration of disease, most patients with type 2 diabetes will eventually fail on oral therapy and become insulin dependent with the necessity for daily injections and multiple daily glucose measurements.

The UKPDS (United Kingdom Prospective Diabetes 25 Study) demonstrated that intensive treatment with metformin, sulfonylureas or insulin resulted in only a limited improvement of glycemic control (difference in HbA1c ~0.9%). In addition, even in patients within the intensive treatment arm glycemic control deteriorated significantly 30 over time and this was attributed to deterioration of β -cell function. Importantly, intensive treatment was not associated with a significant reduction in macrovascular complications, i.e. cardiovascular events. Therefore many patients with type 2 diabetes remain inadequately treated, partly 35 because of limitations in long term efficacy, tolerability and dosing inconvenience of existing antihyperglycemic therapies.

Oral and non-oral antidiabetic drugs conventionally used in therapy (such as e.g. first- or second-line, and/or mono- or 40 121945. (initial or add-on) combination therapy) include, without being restricted thereto, metformin, sulphonylureas, thiazolidinediones, glinides, α -glucosidase inhibitors, GLP-1 or GLP-1 analogues, and insulin or insulin analogues.

The high incidence of therapeutic failure is a major contributor to the high rate of long-term hyperglycemia-associated complications or chronic damages (including micro- and makrovascular complications such as e.g. diabetic nephrophathy, retinopathy or neuropathy, or cardiovascular complications) in patients with type 2 diabetes.

Genetic association studies have identified genetic variations in several genes which are associated with increased risk of type 2 diabetes mellitus. E.g. variations in the genes TCF7L2, KCNJ11 and PPARG independently and interactively increase the risk of progression from impaired fasting 55 glucose and impaired glucose tolerance to overt diabetes. While variation in KCNJ11 may alter insulin secretion and variation in PPARG may alter insulin action, TCF7L2 (transcription factor 7-like 2) is the major susceptibility gene identified to date for type 2 diabetes in various ethnic groups 60 (e.g. Europeans, Indian and Japanese people, Mexican Americans and West Africans). Polymorphisms (single nucleotide polymorphisms, so called SNPs) in TCF7L2, such as e.g. rs12255372 and, particularly, rs7903146, are strongly associated with diabetes. The risk of developing 65 type 2 diabetes is increased by roughly 45% (Odds ratio 1.45) among carriers of one risk T-allele of TCF7L2

4

rs7903146 (CT heterozygotes), and is at least doubled (Odds ratio of 2.41) among TT homozygotes compared to CC homozygotes wild genotypes (Grant et al, Nature Genetics, Vol. 38, 2006, p 320-323). TCF7L2 risk genotypes are associated with increased TCF7L2 expression in pancreatic beta cells, impaired (glucose-stimulated) insulin secretion, incretin effects and enhanced rate of hepatic glucose production as well as predisposition to and prediction of future type 2 diabetes (cf. Lyssenko et al., The Journal of Clinical Investigation, Vol. 117, No 8, 2007, p. 2155-2163). There is evidence that the TCF7L2 rs7903146 risk variants are associated with lower incretin effect on insulin secretion, which may be based, at least in parts, on an impaired sensitivity of the beta cells to incretins.

Thus, diabetes patients harboring TCF7L2 risk variants, particularly carriers of the at risk T-allele of TCF7L2 rs7903146, such as patients harboring the TCF7L2 rs7903146 CT genotype or, particularly, patients harboring the TCF7L2 rs7903146 TT genotype, are expected to be difficult to treat in antidiabetic therapy.

Therefore, there is an unmet medical need for methods, medicaments and pharmaceutical compositions or combinations with a good efficacy with regard to glycemic control, with regard to disease-modifying properties and with regard to reduction of cardiovascular morbidity and mortality while at the same time showing an improved safety profile.

DPP-4 inhibitors represent another novel class of agents that are being developed for the treatment or improvement in glycemic control in patients with type 2 diabetes.

For example, DPP-4 inhibitors and their uses are disclosed in WO 2002/068420, WO 2004/018467, WO 2004/018468, WO 2004/018469, WO 2004/041820, WO 2004/046148, WO 2005/051950, WO 2005/082906, WO 2005/063750, WO 2005/085246, WO 2006/027204, WO 2006/029769, WO2007/014886; WO 2004/050658, WO 2004/11051, WO 2005/058901, WO 2005/097798; WO 2006/068163, WO 2007/071738, WO 2008/017670; WO 2007/128721, WO 2007/128724, WO 2007/128761, or WO 2009/121945.

AIM OF THE PRESENT INVENTION

LP-1 analogues, and insulin or insulin analogues.

The high incidence of therapeutic failure is a major 45 tion and/or method for preventing, slowing progression of, antributor to the high rate of long-term hyperglycemia-sociated complications or chronic damages (including type 2 diabetes mellitus.

A further aim of the present invention is to provide a medication and/or method for improving glycemic control in a patient in need thereof, in particular in patients with type 2 diabetes mellitus, for example in those patients who have variation(s) in one or more genes associated with metabolic diseases (such as e.g. a TCF7L2 risk genotype patient as described herein) or in those patients who are of respective wild-type genotype.

Another aim of the present invention is to provide a medication and/or method for improving glycemic control in a patient with insufficient glycemic control despite monotherapy with an antidiabetic drug, for example metformin, or despite combination therapy with two or three antidiabetic drugs, for example in such a patient who has variation(s) in one or more genes associated with metabolic diseases (such as e.g. a TCF7L2 risk genotype patient as described herein) or in such a patient who is of respective wild-type genotype.

Another aim of the present invention is to provide a medication and/or method for preventing, slowing or delaying progression from impaired glucose tolerance (IGT),

impaired fasting blood glucose (IFG), insulin resistance and/or metabolic syndrome to type 2 diabetes mellitus.

Yet another aim of the present invention is to provide a medication and/or method for preventing, slowing progression of, delaying or treating of a condition or disorder from the group consisting of complications of diabetes mellitus.

A further aim of the present invention is to provide a medication and/or method for reducing the weight or preventing an increase of the weight in a patient in need thereof, for example in such a patient who has variation(s) in one or 10 more genes associated with metabolic diseases (such as e.g. a TCF7L2 risk genotype patient as described herein) or in such a patient who is of respective wild-type genotype.

Another aim of the present invention is to provide a medication with a high efficacy for the treatment of metabolic disorders, in particular of diabetes mellitus, impaired glucose tolerance (IGT), impaired fasting blood glucose (IFG), and/or hyperglycemia, which has good to very good pharmacological and/or pharmacokinetic and/or physicochemical properties. Further aims of the present invention 20 become apparent to the one skilled in the art by description hereinbefore and in the following and by the examples.

SUMMARY OF THE INVENTION

Within the scope of the present invention it has now been found that a DPP-4 inhibitor, preferably linagliptin, as well as a pharmaceutical composition or combination comprising the DPP-4 inhibitor and optionally one or more other active substances (e.g. antidiabetics), is therapeutically effective 30 for improving glycemic control and treating type 2 diabetes mellitus in TCF7L2 rs7903146 CT or TT risk genotype patients and in TCF7L2 rs7903146 CC wild genotype patients.

In particular, it has been found that all investigated 35 TCF7L2 genotype patients (patients with TCF7L2 rs7903146 CT or TT risk genotype or with TCF7L2 rs7903146 CC wild genotype) have a clinically meaningful response to the administered DPP-4 inhibitor, preferably linagliptin.

Thus, within the scope of the present invention, certain subgroups of diabetes patients amenable to antidiabetic therapy according to this invention (comprising using preferably linagliptin, optionally in combination with one or more other active substances such as e.g. other antidiabetics 45 as described herein), include for example, without being limited to, those patients harboring TCF7L2 rs7903146 CC or CT or TT genotype, respectively.

Within the scope of the present invention it has further been found that DPP-4 inhibitors as defined herein as well 50 as pharmaceutical compositions or combinations comprising a DPP-4 inhibitor as defined herein and optionally one or more other active substances can be used in a method of preventing, slowing progression of, delaying (e.g. delaying the onset of) or treating a metabolic disorder (particularly 55 diabetes, especially type 2 diabetes mellitus and conditions related thereto, e.g. diabetic complications), in particular a method for improving glycemic control in a patient, such as in a patient who has variation(s) in one or more genes associated with metabolic diseases (such as e.g. in TCF7L2 60 risk genotype patients as described herein).

Within the scope of the present invention it has further been found that DPP-4 inhibitors as defined herein as well as pharmaceutical compositions or combinations comprising a DPP-4 inhibitor as defined herein and optionally one or 65 more other active substances can be used in a method of preventing, slowing progression of, delaying (e.g. delaying 6

the onset of) or treating a metabolic disorder (particularly diabetes, especially type 2 diabetes mellitus and conditions related thereto), in particular a method for improving glycemic control in a patient, such as in a patient who is of TCF7L2 wild genotype, particularly of TCF7L2 rs7903146 CC wild genotype.

In an embodiment the method comprises the step of identifying a patient being susceptible to the method being used, e.g. comprising testing whether the patient has variation(s) in one or more genes associated with metabolic diseases (e.g. whether the patient is of a TCF7L2 risk genotype as described herein) or whether the patient is of TCF7L2 wild genotype as described herein, and the step of administering such a DPP-4 inhibitor, pharmaceutical composition or combination to the patient determined as being susceptible.

This opens up new therapeutic possibilities in the treatment and prevention of type 2 diabetes mellitus, overweight, obesity, complications of diabetes mellitus and of neighboring disease states, including such patients who have variation(s) in one or more genes associated with metabolic diseases (such as e.g. in TCF7L2 risk genotype patients as described herein) and such patients who are of respective wild-type genotype (such as e.g. TCF7L2 wild genotype patients as described herein).

Moreover, the present invention provides a method for determining of a probability of the likelihood of a favorable response (e.g. in providing glycemic control) or the magnitude of a favorable change in HbA1c of an individual resulting from treating the individual with a DPP-4 inhibitor, preferably linagliptin, or the DPP-4 inhibitor in combination with one or more other active substances (e.g. antidiabetics), said method comprising determining whether the subject is either of TCF7L2 risk genotype (particularly TCF7L2 rs7903146 TT risk genotype) or of TCF7L2 wild genotype (particularly TCF7L2 rs7903146 CC wild genotype), wherein the probability of likelihood of a favorable response 40 or the significantly high magnitude of a favorable change in HbA1c response to administration of the DPP-4 inhibitor. preferably linagliptin, or the DPP-4 inhibitor in combination with one or more other active substances (e.g. antidiabetics)

greater in an individual being of TCF7L2 rs7903146 CC homozygous wild genotype, and lower in an individual of TCF7L2 rs7903146 TT homozygous risk genotype (e.g. but still clinically significant or meaningful).

Therefore, in a one aspect there is provided a pharmaceutical composition or combination comprising

- (a) a DPP-4 inhibitor, and, optionally,
- (b) a second antidiabetic agent selected from the group G3 consisting of biguanides (particularly metformin), thiazolid-indiones, sulfonylureas, glinides, inhibitors of alpha-glucosidase, GLP-1 or GLP-1 analogues and insulin or insulin analogues, and, optionally,
- (c) a third antidiabetic agent being different from (b) selected from the group G3 consisting of biguanides (particularly metformin), thiazolidindiones, sulfonylureas, glinides, inhibitors of alpha-glucosidase, GLP-1 or GLP-1 analogues and insulin or insulin analogues,
- or a pharmaceutically acceptable salt thereof.

In a subaspect there is provided a pharmaceutical composition or combination comprising

- (a) a DPP-4 inhibitor, and, optionally,
- (b) a second antidiabetic agent selected from the group G3 consisting of biguanides (particularly metformin), thiazolid-

indiones, sulfonylureas, glinides, inhibitors of alpha-glucosidase, GLP-1 or GLP-1 analogues and insulin or insulin analogues, and, optionally,

(c) a third antidiabetic agent being different from (b) selected from the group consisting of metformin, a sulfonylurea, 5 pioglitazone, rosiglitazone, repaglinide, nateglinide, acarbose, voglibose, miglitol, GLP-1 or a GLP-1 analogue and insulin or an insulin analogue,

or a pharmaceutically acceptable salt thereof.

In another subaspect there is provided a pharmaceutical 10 composition or combination comprising

(a) a DPP-4 inhibitor, and, optionally,

(b) a second antidiabetic agent selected from the group consisting of metformin, a sulfonylurea, pioglitazone, rosiglitazone, repaglinide, nateglinide, acarbose, voglibose, 15 miglitol, GLP-1 or a GLP-1 analogue and insulin or an insulin analogue, and, optionally,

(c) a third antidiabetic agent being different from (b) selected from the group G3 consisting of biguanides (particularly metformin), thiazolidindiones, sulfonylureas, glinides, 20 inhibitors of alpha-glucosidase, GLP-1 or GLP-1 analogues and insulin or insulin analogues,

or a pharmaceutically acceptable salt thereof.

In a further subaspect there is provided a pharmaceutical composition or combination comprising

(a) a DPP-4 inhibitor, and, optionally,

 (b) a second antidiabetic agent selected from the group consisting of metformin, a sulfonylurea and pioglitazone, and, optionally,

(c) a third antidiabetic agent being different from (b) selected 30 from the group consisting of metformin, a sulfonylurea, pioglitazone, rosiglitazone, repaglinide, nateglinide, acarbose, voglibose, miglitol, GLP-1 or GLP-1 analogue and insulin or insulin analogue,

or a pharmaceutically acceptable salt thereof.

In a further subaspect there is provided a pharmaceutical composition or combination comprising

(a) a DPP-4 inhibitor, and, optionally,

(b) a second antidiabetic agent selected from the group consisting of metformin, a sulfonylurea, pioglitazone, 40 rosiglitazone, repaglinide, nateglinide, acarbose, voglibose, miglitol, GLP-1 or GLP-1 analogue and insulin or insulin analogue, and, optionally,

(c) a third antidiabetic agent being different from (b) selected from the group consisting of metformin, a sulfonylurea and 45 pioglitazone,

or a pharmaceutically acceptable salt thereof.

In a yet further subaspect there is provided a pharmaceutical composition or combination comprising

(a) a DPP-4 inhibitor, and, optionally,

(b) a second antidiabetic agent selected from the group consisting of metformin and pioglitazone, and, optionally, (c) a third antidiabetic agent being different from (b) selected from the group consisting of metformin, a sulfonylurea and pioglitazone.

or a pharmaceutically acceptable salt thereof.

In a yet further subaspect there is provided a pharmaceutical composition or combination comprising

(a) a DPP-4 inhibitor, and, optionally,

(b) a second antidiabetic agent selected from the group 60 consisting of metformin, a sulfonylurea and pioglitazone, and, optionally,

(c) a third antidiabetic agent being different from (b) selected from the group consisting of metformin and pioglitazone, or a pharmaceutically acceptable salt thereof.

When—besides the second anidiabetic agent—a third antidiabetic agent is chosen, said third antidiabetic agent is

8

preferably chosen from another class than the second antidiabetic agent. Thus, it is to be understood that the second and the third antidiabetic agent are different, and preferably they are from different classes (e.g. when the second antidiabetic agent is chosen from the biguanide class, the third antidiabetic agent is preferably chosen from another class). Classes of antidiabetic agents are mentioned above, e.g. biguanide class, thiazolidindione class, sulfonylurea class, glinide class, alpha-glucosidase inhibitor class, GLP-1 analogue class, insulin class, etc.

A particular embodiment of this invention refers to monotherapy with a DPP-4 inhibitor as defined herein and/or to pharmaceutical compositions comprising a DPP-4 inhibitor as sole active ingredient.

Within combinations and/or combination therapy of this invention, a particular embodiment refers to dual combinations and/or dual therapy; another embodiment refers to triple combinations and/or triple therapy.

According to another aspect there is provided a method for preventing, slowing the progression of, delaying or treating a metabolic disorder selected from the group consisting of type 1 diabetes mellitus, type 2 diabetes mellitus, impaired glucose tolerance (IGT), impaired fasting blood glucose (IFG), hyperglycemia, postprandial hyperglycemia, overweight, obesity and metabolic syndrome in a patient in need thereof characterized in that a DPP-4 inhibitor and, optionally, a second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient.

According to another aspect there is provided a method for preventing, slowing the progression of, delaying or treating a metabolic disorder selected from the group consisting of insulin resistance, hyperlipidemia, hypercholesterolemia, dyslipidemia, hypertension, chronic systemic inflammation, retinopathy, neuropathy, nephropathy, atherosclerosis, endothelial dysfunction, non-alcoholic fatty liver disease (NAFLD) and osteoporosis in a patient in need thereof characterized in that a DPP-4 inhibitor and, optionally, a second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient.

According to another aspect there is provided a method for improving glycemic control and/or for reducing of fasting plasma glucose, of postprandial plasma glucose and/or of glycosylated hemoglobin HbA1c in a patient in need thereof characterized in that a DPP-4 inhibitor and, optionally, a second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient.

The pharmaceutical composition of this invention may also have valuable disease-modifying properties with respect to diseases or conditions related to impaired glucose tolerance (IGT), impaired fasting blood glucose (IFG), insu-55 lin resistance and/or metabolic syndrome.

According to another aspect there is provided a method for preventing, slowing, delaying or reversing progression from impaired glucose tolerance (IGT), impaired fasting blood glucose (IFG), insulin resistance and/or from metabolic syndrome to type 2 diabetes mellitus in a patient in need thereof characterized in that a DPP-4 inhibitor and, optionally, a second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient.

As by the use of a pharmaceutical composition or combination of this invention, an improvement of the glycemic control in patients in need thereof is obtainable, also those

conditions and/or diseases related to or caused by an increased blood glucose level may be treated.

According to another aspect there is provided a method for preventing, slowing the progression of, delaying or treating of a condition or disorder selected from the group 5 consisting of complications of diabetes mellitus such as cataracts and micro- and macrovascular diseases, such as nephropathy, retinopathy, neuropathy, learning and memory impairment, neurodegenerative or cognitive disorders, cardio- or cerebrovascular diseases, arteriosclerosis, hyperten- 10 sion, endothelial dysfunction, myocardial infarction, acute coronary syndrome, unstable angina pectoris, stable angina pectoris, cardiomyopathy, heart failure, heart rhythm disorders, vascular restenosis, peripheral arterial occlusive disease, stroke, tissue ischaemia or diabetic foot or ulcus, in a 15 patient in need thereof characterized in that a DPP-4 inhibitor and, optionally, a second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient. In particular one or more aspects of diabetic nephropathy such 20 as hyperperfusion, proteinuria and albuminuria (including micro- or macroalbuminuria) may be treated, their progression slowed or their onset delayed or prevented. The term "tissue ischaemia" particularly comprises diabetic macroangiopathy, diabetic microangiopathy, impaired wound heal- 25 ing and diabetic ulcer. The terms "micro- and macrovascular diseases" and "micro- and macrovascular complications" are used interchangeably in this application.

In an embodiment, by the administration of a pharmaceutical composition or combination of this invention no gain in 30 weight or even a reduction in body weight is the result.

According to another aspect there is provided a method for reducing body weight and/or body fat or preventing an increase in body weight and/or body fat or facilitating a reduction in body weight and/or body fat in a patient in need 35 thereof characterized in that a DPP-4 inhibitor and, optionally, a second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient.

In an embodiment, by an administration of a pharmaceutical composition or combination according to this invention a beta-cell degeneration and a decline of beta-cell functionality such as for example apoptosis or necrosis of pancreatic beta cells can be delayed or prevented. Furthermore, the functionality of pancreatic cells can be improved or restored, 45 and the number and size of pancreatic beta cells increased. It may be shown that the differentiation status and hyperplasia of pancreatic beta-cells disturbed by hyperglycemia can be normalized by treatment with a pharmaceutical composition or combination of this invention.

According to another aspect there is provided a method for preventing, slowing, delaying or treating the degeneration of pancreatic beta cells and/or the decline of the functionality of pancreatic beta cells and/or for improving and/or restoring the functionality of pancreatic beta cells second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient.

In an embodiment, by the administration of a pharmaceutical composition or combination of the present invention, an abnormal accumulation of (ectopic) fat, in particular in the liver, may be reduced or inhibited.

According to another aspect there is provided a method 65 for preventing, slowing, delaying or treating diseases or conditions attributed to an abnormal accumulation of liver or

10

ectopic fat in a patient in need thereof characterized in that a DPP-4 inhibitor and, optionally, a second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient. Diseases or conditions which are attributed to an abnormal accumulation of liver or ectopic fat are particularly selected from the group consisting of general fatty liver, non-alcoholic fatty liver (NAFL), non-alcoholic steatohepatitis (NASH), hyperalimentation-induced fatty liver, diabetic fatty liver, alcoholic-induced fatty liver or toxic fatty liver, particularly non-alcoholic fatty liver disease (NAFLD), including hepatic steatosis, non-alcoholic steatohepatitis (NASH) and/or liver fibrosis.

According to a further aspect of the present invention, there is provided a method for preventing, slowing the progression, delaying, attenuating, treating or reversing hepatic steatosis, (hepatic) inflammation and/or an abnormal accumulation of liver fat in a patient in need thereof characterized in that a DPP-4 inhibitor and, optionally, a second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient.

According to another aspect there is provided a method for maintaining and/or improving the insulin sensitivity and/or for treating or preventing hyperinsulinemia and/or insulin resistance in a patient in need thereof characterized in that a DPP-4 inhibitor and, optionally, a second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient.

According to another aspect of the invention, there is provided a method for preventing, slowing progression of, delaying, or treating new onset diabetes after transplantation (NODAT) and/or post-transplant metabolic syndrome (PTMS) in a patient in need thereof characterized in that a DPP-4 inhibitor and, optionally, a second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient

According to a further aspect of the invention, there is provided a method for preventing, delaying, or reducing NODAT and/or PTMS associated complications including micro- and macrovascular diseases and events, graft rejection, infection, and death in a patient in need thereof characterized in that a DPP-4 inhibitor and, optionally, a second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient.

According to another aspect of the invention, there is provided a method for treating hyperuricemia and hyperuricemia-associated conditions, such as for example gout, hypertension and renal failure, in a patient in need thereof characterized in that a DPP-4 inhibitor and, optionally, a second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient.

According to another aspect there is provided the use of a DPP-4 inhibitor for the manufacture of a medicament for use in a method of

preventing, slowing the progression of, delaying or treating a metabolic disorder selected from the group consisting of type 1 diabetes mellitus, type 2 diabetes mellitus, impaired glucose tolerance (IGT), impaired fasting blood glucose (IFG), hyperglycemia, postprandial hyperglycemia, overweight, obesity and metabolic syndrome; or

improving glycemic control and/or for reducing of fasting plasma glucose, of postprandial plasma glucose and/or of glycosylated hemoglobin HbA1c; or

preventing, slowing, delaying or reversing progression from impaired glucose tolerance (IGT), impaired fasting blood glucose (IFG), insulin resistance and/or from metabolic syndrome to type 2 diabetes mellitus; or

preventing, slowing the progression of, delaying or treating of a condition or disorder selected from the group consisting of complications of diabetes mellitus such as cataracts and micro- and macrovascular diseases, such as nephropathy, retinopathy, neuropathy, tissue ischaemia, arteriosclerosis, myocardial infarction, stroke and peripheral arterial occlusive disease; or

reducing body weight and/or body fat or preventing an 15 increase in body weight and/or body fat or facilitating a reduction in body weight and/or body fat; or

preventing, slowing, delaying or treating the degeneration of pancreatic beta cells and/or the decline of the functionality of pancreatic beta cells and/or for improving 20 and/or restoring or protecting the functionality of pancreatic beta cells and/or restoring the functionality of pancreatic insulin secretion; or

preventing, slowing, delaying or treating diseases or conditions attributed to an abnormal accumulation of liver 25 or ectopic fat; or

maintaining and/or improving the insulin sensitivity and/ or for treating or preventing hyperinsulinemia and/or insulin resistance; or

preventing, slowing progression of, delaying, or treating 30 new onset diabetes after transplantation (NODAT) and/ or post-transplant metabolic syndrome (PTMS); or

preventing, delaying, or reducing NODAT and/or PTMS associated complications including micro- and macro-vascular diseases and events, graft rejection, infection, 35 and death: or

treating hyperuricemia and hyperuricemia associated conditions:

in a patient in need thereof, comprising administering the DPP-4 inhibitor alone or, optionally, in combination with a 40 second and, optionally, with a third antidiabetic agent as defined hereinbefore and hereinafter to the patient.

According to another aspect there is provided the use of a second antidiabetic agent as defined hereinbefore and hereinafter for the manufacture of a medicament for use in 45 a method of

preventing, slowing the progression of, delaying or treating a metabolic disorder selected from the group consisting of type 1 diabetes mellitus, type 2 diabetes mellitus, impaired glucose tolerance (IGT), impaired 50 fasting blood glucose (IFG), hyperglycemia, postprandial hyperglycemia, overweight, obesity and metabolic syndrome: or

improving glycemic control and/or for reducing of fasting plasma glucose, of postprandial plasma glucose and/or 55 of glycosylated hemoglobin HbA1c; or

preventing, slowing, delaying or reversing progression from impaired glucose tolerance (IGT), impaired fasting blood glucose (IFG), insulin resistance and/or from metabolic syndrome to type 2 diabetes mellitus; or

preventing, slowing the progression of, delaying or treating of a condition or disorder selected from the group consisting of complications of diabetes mellitus such as cataracts and micro- and macrovascular diseases, such as nephropathy, retinopathy, neuropathy, tissue ischaemia, arteriosclerosis, myocardial infarction, stroke and peripheral arterial occlusive disease; or

12

reducing body weight and/or body fat or preventing an increase in body weight and/or body fat or facilitating a reduction in body weight and/or body fat; or

preventing, slowing, delaying or treating the degeneration of pancreatic beta cells and/or the decline of the functionality of pancreatic beta cells and/or for improving and/or restoring the functionality of pancreatic beta cells and/or restoring the functionality of pancreatic insulin secretion; or

preventing, slowing, delaying or treating diseases or conditions attributed to an abnormal accumulation of liver or ectopic fat; or

maintaining and/or improving the insulin sensitivity and/ or for treating or preventing hyperinsulinemia and/or insulin resistance;

in a patient in need thereof, comprising administering the second antidiabetic agent in combination with a DPP-4 inhibitor and, optionally, with a third antidiabetic agent as defined hereinbefore and hereinafter to the patient.

According to another aspect there is provided the use of a pharmaceutical composition according to the present invention for the manufacture of a medicament for a therapeutic and preventive method as described hereinbefore and hereinafter.

Patients of a TCF7L2 risk genotype (also referred to herein as TCF7L2 risk genotype patients) within the meaning of this invention refer to those patients who have one or more single nucleotide polymorphisms (SNPs) in the gene coding for TCF7L2, especially a SNP selected from rs7903146, rs12255372 and rs10885406, especially rs7903146; in more particular, those patients who carry at least one T allele of SNP rs7903146 of TCF7L2, i.e. the CT genotype or TT genotype; especially those who carry two T alleles of SNP rs7903146 of TCF7L2, i.e. the TT genotype, are at high-risk and are expected to be difficult to treat (e.g. to achieve adequate glycemic control).

The present invention provides a DPP-4 inhibitor (preferably linagliptin), pharmaceutical composition, combination or medicament according to the present invention for use in a therapeutic and/or preventive method as described hereinbefore and hereinafter (e.g. treating type 2 diabetes) in one or more of the following patient groups:

TCF7L2 high risk genotype patients carrying two T alleles of SNP rs7903146 of TCF7L2, i.e. TT genotype (where clinically meaningful response e.g. in glycemic control is provided),

TCF7L2 risk genotype patients carrying one T allele of SNP rs7903146 of TCF7L2, i.e. CT genotype (where clinically favorable response e.g. in glycemic control is provided),

TCF7L2 wild genotype patients carrying two CC alleles of SNP rs7903146 of TCF7L2, i.e. CC genotype (where clinically more favorable response e.g. in glycemic control is provided).

Within a particular aspect of the invention, the invention relates to a DPP-4 inhibitor, a pharmaceutical composition or combination of the present invention for a therapeutic and/or preventive method or use as described hereinbefore and hereinafter (e.g. treating type 2 diabetes), said method or use comprising

(i) identifying a patient being susceptible to said therapeutic and/or preventive method or use comprising testing whether the patient is of any TCF7L2 risk genotype, particularly whether the patient has one or more single nucleotide polymorphisms (SNPs) in the gene coding for TCF7L2, especially a SNP selected from rs7903146, rs12255372 and rs10885406, for

example whether the patient carries at least one T allele of SNP rs7903146 of TCF7L2, e.g. whether the patient is of CT genotype (i.e. whether the patient carries one T allele of SNP rs7903146 of TCF7L2) or, particularly, whether the patient is of TT genotype (i.e. whether the 5 patient carries two T alleles of SNP rs7903146 of TCF7L2), or testing whether the patient is of TCF7L2 wild genotype, particularly whether the patient carries two C alleles of SNP rs7903146 of TCF7L2 (i.e. whether the patient is of CC wild genotype), and

(ii) administering an effective amount of the DPP-4 inhibitor, pharmaceutical composition or combination to the patient identified in step (i).

Within another particular aspect of the invention, the invention relates to a DPP-4 inhibitor, a pharmaceutical 15 composition, combination or medicament of the present invention for a therapeutic and/or preventive method or use as described hereinbefore and hereinafter (e.g. treating type 2 diabetes) in TCF7L2 risk genotype patients, e.g. in those patients who have one or more single nucleotide polymor- 20 phisms (SNPs) in the gene coding for TCF7L2, especially a SNP selected from rs7903146, rs12255372 and rs10885406, especially rs7903146; in more particular, in those patients who carry at least one T allele of SNP rs7903146 of TCF7L2, i.e. the CT genotype or TT genotype.

Within another particular aspect of the invention, the invention relates to a DPP-4 inhibitor, a pharmaceutical composition, combination or medicament of the present invention for a therapeutic and/or preventive method or use as described hereinbefore and hereinafter (e.g. treating type 30 2 diabetes) in TCF7L2 wild genotype patients, e.g. in those patients who carry two C alleles of SNP rs7903146 of TCF7L2, i.e. the CC genotype.

In this context, a particular sub-population of the patients described hereinbefore and hereinafter (e.g. of the patients in 35 need of a therapeutic or preventive method as described herein), refers to those patients who have one or more single nucleotide polymorphisms (SNPs) in the gene coding for TCF7L2, especially at least one SNP selected from rs7903146, rs12255372 and rs10885406, especially 40 rs7903146, in more particular, those patients who carry at least one T allele of SNP rs7903146 of TCF7L2, i.e. the CT genotype or TT genotype.

In more particular, those patients who carry at least one T allele of SNP rs7903146 of TCF7L2, i.e. the CT genotype or 45 TT genotype, especially who carry two T alleles of SNP rs7903146 of TCF7L2, i.e. the TT genotype, are strongly susceptible to increased TCF7L2 expression in pancreatic beta cells, impaired insulin secretion, incretine effects, enhanced rate of hepatic glucose production and/or diabetes. 50 The T allele of rs7903146 TCF7L2 is associated with impaired insulinotropic action of incretin hormones, reduced 24 h profiles of plasma insulin and glucagon, and increased hepatic glucose production.

Another particular sub-population of the patients 55 described hereinbefore and hereinafter (e.g. of the patients in need of a therapeutic or preventive method as described herein), refers to those patients who are of TCF7L2 wild genotype, particularly those who are of the TCF7L2 rs7903146 CC wild genotype.

According to one embodiment of this aspect of the invention, there is provided a DPP-4 inhibitor, a pharmaceutical composition, combination or medicament according to the present invention for a therapeutic and/or preventive method or use as described hereinbefore and hereinafter 65 (particularly for treating and/or preventing type 2 diabetes and/or obesity), in patients with reduced (glucose-stimu14

lated) insulin secretion, increased hepatic gluconeogenesis and/or reduced insulinotropic effect or action of incretin hormones (e.g. GLP-1 and/or GIP), e.g. impaired incretin sensitivity, associated with a TCF7L2 risk genotype, particularly with such a TCF7L2 risk genotype as mentioned

According to another embodiment of this aspect of the invention, there is provided a method of determining patient's treatment response to a DPP-4 inhibitor, a pharmaceutical composition, combination or medicament according to the present invention, said method comprising the step of determining whether the patient is of TCF7L2 risk genotype as described herein, e.g. testing whether the patient belongs to the particular subpopulation of TCF7L2 risk genotype carriers, or determining whether the patient is of TCF7L2 wild genotype, e.g. testing whether the patient carries the wild-type CC allele at rs7903146 in TCF7L2.

According to another embodiment of this aspect of the invention, there is provided a DPP-4 inhibitor, a pharmaceutical composition, combination or medicament according to the present invention for use in a therapeutic and/or preventive method as described hereinbefore and hereinafter (particularly for treating and/or preventing type 2 diabetes and/or obesity) in a patient in need thereof, said method comprising testing whether the patient is of any TCF7L2 risk genotype as described herein.

According to another embodiment of this aspect of the invention, there is provided a DPP-4 inhibitor, a pharmaceutical composition, combination or medicament according to the present invention for use in a therapeutic and/or preventive method as described hereinbefore and hereinafter (particularly for treating and/or preventing type 2 diabetes and/or obesity) in a patient in need thereof, said method comprising testing whether the patient is of TCF7L2 wild genotype as described herein.

According to another aspect of the invention, the testing for TCF7L2 risk genotypes may be used for patient stratification, e.g. to enrich patient population in clinical trials to test the efficacy of the DPP-4 inhibitor.

According to another aspect of the invention, the method of determining the treatment susceptibility of an individual (e.g. comprising the testing for TCF7L2 risk or wild genotypes as described herein) may be used for determination whether the patient may respond to a lower level or may require a higher level of administered DPP-4 inhibitor, optionally in combination with one or more other active substances.

According to another aspect of the invention, determining the treatment susceptibility of an individual comprising the testing for TCF7L2 risk or wild genotypes as described herein may be used for determination whether the patient may be treated in monotherapy or in combination therapy with one or more additional antidiabetics according to this invention, e.g. to provide adequate glycemic control. For example, those patients with decreased likelihood of favorable response may require combination treatment, e.g. to achieve adequate glycemic control.

DEFINITIONS

60

The term "active ingredient" of a pharmaceutical composition or combination of the present invention means the DPP-4 inhibitor and/or, if present, the second antidiabetic agent and/or, if present, the third antidiabetic agent of the present invention.

The term "body mass index" or "BMI" of a human patient is defined as the weight in kilograms divided by the square of the height in meters, such that BMI has units of kg/m².

The term "overweight" is defined as the condition wherein the individual has a BMI greater than or 25 kg/m² and less than 30 kg/m². The terms "overweight" and "preobese" are used interchangeably.

The term "obesity" is defined as the condition wherein the individual has a BMI equal to or greater than 30 kg/m². According to a WHO definition the term obesity may be categorized as follows: the term "class I obesity" is the condition wherein the BMI is equal to or greater than 30 kg/m² but lower than 35 kg/m²; the term "class II obesity" is the condition wherein the BMI is equal to or greater than 35 kg/m² but lower than 40 kg/m²; the term "class III obesity" is the condition wherein the BMI is equal to or greater than 40 kg/m².

The term "visceral obesity" is defined as the condition wherein a waist-to-hip ratio of greater than or equal to 1.0_{20} in men and 0.8 in women is measured. It defines the risk for insulin resistance and the development of pre-diabetes.

The term "abdominal obesity" is usually defined as the condition wherein the waist circumference is >40 inches or 102 cm in men, and is >35 inches or 94 cm in women. With 25 regard to a Japanese ethnicity or Japanese patients abdominal obesity may be defined as waist circumference ≥85 cm in men and ≥90 cm in women (see e.g. investigating committee for the diagnosis of metabolic syndrome in Japan).

The term "euglycemia" is defined as the condition in which a subject has a fasting blood glucose concentration within the normal range, greater than 70 mg/dL (3.89 mmol/L) and less than 110 mg/dL (6.11 mmol/L) or 100 mg mg/dL (5.6 mmol/L). The word "fasting" has the usual meaning as a medical term.

The term "hyperglycemia" is defined as the condition in which a subject has a fasting blood glucose concentration above the normal range, greater than 110 mg/dL (6.11 $_{
m 40}$ mmol/L) or 100 mg mg/dL (5.6 mmol/L). The word "fasting" has the usual meaning as a medical term.

The term "hypoglycemia" is defined as the condition in which a subject has a blood glucose concentration below the normal range of 60 to 115 mg/dL (3.3 to 6.3 mmol/L), in 45 particular below 70 mg/dL (3.89 mmol/L).

The term "postprandial hyperglycemia" is defined as the condition in which a subject has a 2 hour postprandial blood glucose or serum glucose concentration greater than 200 mg/dL (11.11 mmol/L).

The term "impaired fasting blood glucose" or "IFG" is defined as the condition in which a subject has a fasting blood glucose concentration or fasting serum glucose concentration in a range from 100 to 125 mg/dl (i.e. from 5.6 to 6.9 mmol/l), in particular greater than 110 mg/dL and less 55 than 126 mg/dl (7.00 mmol/L). A subject with "normal fasting glucose" has a fasting glucose concentration smaller than 100 mg/dl, i.e. smaller than 5.6 mmol/l.

The term "impaired glucose tolerance" or "IGT" is defined as the condition in which a subject has a 2 hour 60 postprandial blood glucose or serum glucose concentration greater than 140 mg/dl (7.78 mmol/L) and less than 200 mg/dL (11.11 mmol/L). The abnormal glucose tolerance, i.e. the 2 hour postprandial blood glucose or serum glucose concentration can be measured as the blood sugar level in 65 mg of glucose per dL of plasma 2 hours after taking 75 g of glucose after a fast. A subject with "normal glucose toler-

16

ance" has a 2 hour postprandial blood glucose or serum glucose concentration smaller than 140 mg/dl (7.78 mmol/ I.)

The term "hyperinsulinemia" is defined as the condition in which a subject with insulin resistance, with or without euglycemia, has fasting or postprandial serum or plasma insulin concentration elevated above that of normal, lean individuals without insulin resistance, having a waist-to-hip ratio <1.0 (for men) or <0.8 (for women).

The terms "insulin-sensitizing", "insulin resistance-improving" or "insulin resistance-lowering" are synonymous and used interchangeably.

The term "insulin resistance" is defined as a state in which circulating insulin levels in excess of the normal response to a glucose load are required to maintain the euglycemic state (Ford E S, et al. JAMA. (2002) 287:356-9). A method of determining insulin resistance is the euglycaemic-hyperinsulinaemic clamp test. The ratio of insulin to glucose is determined within the scope of a combined insulin-glucose infusion technique. There is found to be insulin resistance if the glucose absorption is below the 25th percentile of the background population investigated (WHO definition). Rather less laborious than the clamp test are so called minimal models in which, during an intravenous glucose tolerance test, the insulin and glucose concentrations in the blood are measured at fixed time intervals and from these the insulin resistance is calculated. With this method, it is not possible to distinguish between hepatic and peripheral insulin resistance.

Furthermore, insulin resistance, the response of a patient with insulin resistance to therapy, insulin sensitivity and hyperinsulinemia may be quantified by assessing the "homeostasis model assessment to insulin resistance (HOMA-IR)" score, a reliable indicator of insulin resistance (Katsuki A, et al. Diabetes Care 2001; 24: 362-5). Further reference is made to methods for the determination of the HOMA-index for insulin sensitivity (Matthews et al., Diabetologia 1985, 28:412-19), of the ratio of intact proinsulin to insulin (Forst et al., Diabetes 2003, 52(Suppl. 1): A459) and to an euglycemic clamp study. In addition, plasma adiponectin levels can be monitored as a potential surrogate of insulin sensitivity. The estimate of insulin resistance by the homeostasis assessment model (HOMA)-IR score is calculated with the formula (Galvin P, et al. Diabet Med 1992; 9:921-8):

HOMA-IR=[fasting serum insulin (μU/mL)]x[fasting plasma glucose (mmol/L)/22.5]

As a rule, other parameters are used in everyday clinical practice to assess insulin resistance. Preferably, the patient's triglyceride concentration is used, for example, as increased triglyceride levels correlate significantly with the presence of insulin resistance.

Patients with a predisposition for the development of IGT or IFG or type 2 diabetes are those having euglycemia with hyperinsulinemia and are by definition, insulin resistant. A typical patient with insulin resistance is usually overweight or obese. If insulin resistance can be detected, this is a particularly strong indication of the presence of pre-diabetes. Thus, it may be that in order to maintain glucose homoeostasis a person needs 2-3 times as much insulin as a healthy person, without this resulting in any clinical symptoms

The methods to investigate the function of pancreatic beta-cells are similar to the above methods with regard to insulin sensitivity, hyperinsulinemia or insulin resistance: An improvement of beta-cell function can be measured for

example by determining a HOMA-index for beta-cell function (Matthews et al., Diabetologia 1985, 28:412-19), the ratio of intact proinsulin to insulin (Forst et al., Diabetes 2003, 52(Suppl. 1): A459), the insulin/C-peptide secretion after an oral glucose tolerance test or a meal tolerance test, 5 or by employing a hyperglycemic clamp study and/or minimal modeling after a frequently sampled intravenous glucose tolerance test (Stumvoll et al., Eur J Clin Invest 2001, 31: 380-81).

The term "pre-diabetes" is the condition wherein an 10 individual is pre-disposed to the development of type 2 diabetes. Pre-diabetes extends the definition of impaired glucose tolerance to include individuals with a fasting blood glucose within the high normal range 100 mg/dL (J. B. Meigs, et al. Diabetes 2003; 52:1475-1484) and fasting 15 hyperinsulinemia (elevated plasma insulin concentration). The scientific and medical basis for identifying pre-diabetes as a serious health threat is laid out in a Position Statement entitled "The Prevention or Delay of Type 2 Diabetes" issued jointly by the American Diabetes Association and the 20 National Institute of Diabetes and Digestive and Kidney Diseases (Diabetes Care 2002; 25:742-749).

Individuals likely to have insulin resistance are those who have two or more of the following attributes: 1) overweight or obese, 2) high blood pressure, 3) hyperlipidemia, 4) one 25 or more 1st degree relative with a diagnosis of IGT or IFG or type 2 diabetes. Insulin resistance can be confirmed in these individuals by calculating the HOMA-IR score. For the purpose of this invention, insulin resistance is defined as the clinical condition in which an individual has a HOMA- 30 IR score >4.0 or a HOMA-IR score above the upper limit of normal as defined for the laboratory performing the glucose and insulin assays.

The term "type 2 diabetes" is defined as the condition in which a subject has a fasting blood glucose or serum glucose 35 E, et al. Am J. Epidemiol. (2002) 156:1070-7). Triglycerides concentration greater than 125 mg/dL (6.94 mmol/L). The measurement of blood glucose values is a standard procedure in routine medical analysis. If a glucose tolerance test is carried out, the blood sugar level of a diabetic will be in excess of 200 mg of glucose per dL (11.1 mmol/l) of plasma 40 2 hours after 75 g of glucose have been taken on an empty stomach. In a glucose tolerance test 75 g of glucose are administered orally to the patient being tested after 10-12 hours of fasting and the blood sugar level is recorded immediately before taking the glucose and 1 and 2 hours 45 after taking it. In a healthy subject, the blood sugar level before taking the glucose will be between 60 and 110 mg per dL of plasma, less than 200 mg per dL 1 hour after taking the glucose and less than 140 mg per dL after 2 hours. If after 2 hours the value is between 140 and 200 mg, this is 50 regarded as abnormal glucose tolerance.

The term "late stage type 2 diabetes mellitus" includes type 2 diabetes patients with a secondary antidiabetic drug failure, indication for insulin therapy and progression to micro- and macrovascular complications e.g. diabetic neph- 55 ropathy, or coronary heart disease (CHD).

The term "HbA1c" refers to the product of a non-enzymatic glycation of the haemoglobin B chain. Its determination is well known to one skilled in the art. In monitoring the treatment of diabetes mellitus the HbA1c value is of excep- 60 tional importance. As its production depends essentially on the blood sugar level and the life of the erythrocytes, the HbA1c in the sense of a "blood sugar memory" reflects the average blood sugar levels of the preceding 4-6 weeks. Diabetic patients whose HbA1c value is consistently well 65 adjusted by intensive diabetes treatment (i.e. <6.5% of the total haemoglobin in the sample), are significantly better

18

protected against diabetic microangiopathy. For example, metformin on its own achieves an average improvement in the HbA1c value in the diabetic of the order of 1.0-1.5%. This reduction of the HbA1C value is not sufficient in all diabetics to achieve the desired target range of <6.5% and preferably <6% HbA1c.

The term "insufficient glycemic control" or "inadequate glycemic control" in the scope of the present invention means a condition wherein patients show HbA1c values above 6.5%, in particular above 7.0%, even more preferably above 7.5%, especially above 8%.

The "metabolic syndrome", also called "syndrome X" (when used in the context of a metabolic disorder), also called the "dysmetabolic syndrome" is a syndrome complex with the cardinal feature being insulin resistance (Laaksonen DE, et al. Am J Epidemiol 2002; 156:1070-7). According to the ATP III/NCEP guidelines (Executive Summary of the Third Report of the National Cholesterol Education Program (NCEP) Expert Panel on Detection, Evaluation, and Treatment of High Blood Cholesterol in Adults (Adult Treatment Panel III) JAMA: Journal of the American Medical Association (2001) 285:2486-2497), diagnosis of the metabolic syndrome is made when three or more of the following risk factors are present:

- 1. Abdominal obesity, defined as waist circumference >40 inches or 102 cm in men, and >35 inches or 94 cm in women; or with regard to a Japanese ethnicity or Japanese patients defined as waist circumference ≥85 cm in men and ≥90 cm in women;
- 2. Triglycerides: ≥150 mg/dL
- 3. HDL-cholesterol <40 mg/dL in men
- 4. Blood pressure ≥130/85 mm Hg (SBP ≥130 or DBP
- 5. Fasting blood glucose ≥110 mg/dL or ≥100 mg/dL

The NCEP definitions have been validated (Laaksonen D and HDL cholesterol in the blood can also be determined by standard methods in medical analysis and are described for example in Thomas L (Editor): "Labor and Diagnose", TH-Books Verlagsgesellschaft mbH, Frankfurt/Main, 2000.

According to a commonly used definition, hypertension is diagnosed if the systolic blood pressure (SBP) exceeds a value of 140 mm Hg and diastolic blood pressure (DBP) exceeds a value of 90 mm Hg. If a patient is suffering from manifest diabetes it is currently recommended that the systolic blood pressure be reduced to a level below 130 mm Hg and the diastolic blood pressure be lowered to below 80 mm Hg.

The definitions of NODAT (new onset diabetes after transplantation) and PTMS (post-transplant metabolic syndrome) follow closely that of the American Diabetes Association diagnostic criteria for type 2 diabetes, and that of the International Diabetes Federation (IDF) and the American Heart Association/National Heart, Lung, and Blood Institute, for the metabolic syndrome. NODAT and/or PTMS are associated with an increased risk of micro- and macrovascular disease and events, graft rejection, infection, and death. A number of predictors have been identified as potential risk factors related to NODAT and/or PTMS including a higher age at transplant, male gender, the pretransplant body mass index, pre-transplant diabetes, and immunosuppression.

The term "hyperuricemia" denotes a condition of high serum total urate levels. In human blood, uric acid concentrations between 3.6 mg/dL (ca. 214 µmol/L) and 8.3 mg/dL (ca. 494 µmol/L) are considered normal by the American Medical Association. High serum total urate levels, or hyperuricemia, are often associated with several maladies.

For example, high serum total urate levels can lead to a type of arthritis in the joints known as gout. Gout is a condition created by a build up of monosodium urate or uric acid crystals on the articular cartilage of joints, tendons and surrounding tissues due to elevated concentrations of total urate levels in the blood stream. The build up of urate or uric acid on these tissues provokes an inflammatory reaction of these tissues. Saturation levels of uric acid in urine may result in kidney stone formation when the uric acid or urate crystallizes in the kidney. Additionally, high serum total urate levels are often associated with the so-called metabolic syndrome, including cardiovascular disease and hypertension

The term "DPP-4 inhibitor" in the scope of the present 15 invention relates to a compound that exhibits inhibitory activity on the enzyme dipeptidyl peptidase IV (DPP-4). Such inhibitory activity can be characterised by the IC50 value. A DPP-4 inhibitor preferably exhibits an IC50 value 20 below 10000 nM, preferably below 1000 nM. Certain DPP-4 inhibitors exhibit an IC50 value below 100 nM, or even ≤50 nM. IC50 values of DPP-4 inhibitors are usually above 0.01 nM, or even above 0.1 nM. DPP-IV inhibitors may include 25 biologic and non-biologic, in particular non-peptidic compounds. The inhibitory effect on DPP-4 can be determined by methods known in the literature, in particular as described in the application WO 02/068420 or WO 2004/ 30 018468 (page 34), which are incorporated herein by reference in its entirety. The term "DPP-4 inhibitor" also comprises any pharmaceutically acceptable salts thereof, hydrates and solvates thereof, including the respective crys- 35 talline forms.

The terms "treatment" and "treating" or analogous terms comprise particularly therapeutic treatment of patients having already developed said condition, in particular in manifest form. Therapeutic treatment may be symptomatic treatment in order to relieve the symptoms of the specific indication or causal treatment in order to reverse or partially reverse the conditions of the indication or to stop or slow down progression of the disease. Thus the compositions and methods of the present invention may be used for instance as therapeutic treatment over a period of time as well as for chronic therapy.

The terms "prophylactically treating", "preventive treating" and "preventing" or ananlogous terms are used interchangeably and comprise a treatment of patients at risk to develop a condition mentioned hereinbefore, thus reducing said risk.

DETAILED DESCRIPTION

The aspects of the present invention, in particular the pharmaceutical compounds, compositions, combinations, methods and uses, refer to DPP-4 inhibitors, second and/or third antidiabetic agents as defined hereinbefore and hereinafter.

In a first embodiment (embodiment A), a DPP-4 inhibitor 65 in the context of the present invention is any DPP-4 inhibitor of

formula (I)

$$R1$$
N
N
R2

or

formula (II)

$$\begin{array}{c} & \text{(II)} \\ & \text{N} \\ & \text{N} \\ & \text{N} \\ & \text{N} \end{array}$$

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

wherein R1 denotes ([1,5]naphthyridin-2-yl)methyl, (quinazolin-2-yl)methyl, (quinoxalin-6-yl)methyl, (4-methyl-quinazolin-2-yl)methyl, 2-cyano-benzyl, (3-cyano-quinolin-2-yl)methyl, (3-cyano-pyridin-2-yl)methyl, (4-methyl-pyrimidin-2-yl)methyl, or (4,6-dimethyl-pyrimidin-2-yl)methyl and R2 denotes 3-(R)-amino-piperidin-1-yl, (2-amino-2-methyl-propyl)-methylamino or (2-(S)-amino-propyl)-methylamino, or its pharmaceutically acceptable salt.

In a second embodiment (embodiment B), a DPP-4 inhibitor in the context of the present invention is a DPP-4 inhibitor selected from the group consisting of sitagliptin, vildagliptin, saxagliptin, alogliptin, gemigliptin,

(2S)-1-{[2-(5-Methyl-2-phenyl-oxazol-4-yl)-ethylamino]-acetyl}-pyrrolidine-2-carbonitrile,

55 (2S)-1-{[1,1,-Dimethyl-3-(4-pyridin-3-yl-imidazol-1-yl)-propylamino]-acetyl}-pyrrolidine-2-carbonitrile,

(S)-1-((2S,3S,11bS)-2-Amino-9,10-dimethoxy-1,3,4,6,7, 11b-hexahydro-2H-pyrido[2,1-a]isoquinolin-3-yl)-4-fluoromethyl-pyrrolidin-2-one,

(3,3-Difluoropyrrolidin-1-yl)-((2S,4S)-4-(4-(pyrimidin-2-yl)piperazin-1-yl)pyrrolidin-2-yl)methanone,

(1((3S,4S)-4-amino-1-(4-(3,3-difluoropyrrolidin-1-yl)-1,3, 5-triazin-2-yl)pyrrolidin-3-yl)-5,5-difluoropiperidin-2-one.

5 (2S,4S)-1-{2-[(3S,1R)-3-(1H-1,2,4-Triazol-1-ylmethyl)cyclopentylamino]-acetyl}-4-fluoropyrrolidine-2-carbonitrile,

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(R)-2-[6-(3-Amino-piperidin-1-yl)-3-methyl-2,4-dioxo-3,4-dihydro-2H-pyrimidin-1-ylmethyl]-4-fluoro-benzonitrile,

5-{(S)-2-[2-((S)-2-Cyano-pyrrolidin-1-yl)-2-oxo-ethyl-amino]-propyl}-5-(1H-tetrazol-5-yl)-10,11-dihydro-5H-dibenzo[a,d]cycloheptene-2,8-dicarboxylic acid bis-dimethylamide,

3-{(2S,4S)-4-[4-(3-Methyl-1-phenyl-1H-pyrazol-5-yl)pip-erazin-1-yl]pyrrolidin-2-ylcarbonyl}thiazolidine,

[(2R)-1-{[(3R)-pyrrolidin-3-ylamino]acetyl}pyrrolidin-2-yl]boronic acid,

(2\$,4\$)-1-[2-[(4-ethoxycarbonylbicyclo[2.2.2]oct-1-yl) amino]acetyl]-4-fluoropyrrolidine-2-carbonitrile,

2-({6-[(3R)-3-amino-3-methylpiperidin-1-yl]-1,3-dimethyl-2,4-dioxo-1,2,3,4-tetrahydro-5H-pyrrolo[3,2-d]pyrimi-din-5-yl}methyl)-4-fluorobenzonitrile,

6-[(3R)-3-amino-piperidin-1-yl]-5-(2-chloro-5-fluoro-benzyl)-1,3-dimethyl-1,5-dihydro-pyrrolo[3,2-d]pyrimidine-2,4-dione, and

(S)-2-methylpyrazolo[1,5-a]primidine-6-carboxylic acid{2-[(2-cyanopyrrolidin-1-yl)-2-oxoethylamino]-2-methylpropyl}amide,

or its pharmaceutically acceptable salt.

Regarding the first embodiment (embodiment A), preferred DPP-4 inhibitors are any or all of the following compounds and their pharmaceutically acceptable salts:

1-[(4-methyl-quinazolin-2-yl)methyl]-3-methyl-7-(2-butyn-1-yl)-8-(3-(R)-amino-piperidin-1-yl)-xanthine (compare WO 2004/018468, example 2 (142)):

$$\begin{array}{c|c}
 & 30 \\
 & N \\
 & N$$

1-[([1,5]naphthyridin-2-yl)methyl]-3-methyl-7-(2-butyn-1-yl)-8-((R)-3-amino-piperidin-1-yl)-xanthine (compare WO 2004/018468, example 2 (252)):

1-[(Quinazolin-2-yl)methyl]-3-methyl-7-(2-butyn-1-yl)-8-((R)-3-amino-piperidin-1-yl)-xanthine (compare WO 2004/018468, example 2 (80)):

2-((R)-3-Amino-piperidin-1-yl)-3-(but-2-yinyl)-5-(4-methyl-quinazolin-2-ylmethyl)-3,5-dihydro-imidazo [4,5-d]pyridazin-4-one (compare WO 2004/050658, example 136):

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ &$$

1-[(4-Methyl-quinazolin-2-yl)methyl]-3-methyl-7-(2-butyin-1-yl)-8-[(2-amino-2-methyl-propyl)-methylamino]-xanthine (compare WO 2006/029769, example 2 (1)):

1-[(3-Cyano-quinolin-2-yl)methyl]-3-methyl-7-(2-butyn-1-yl)-8-((R)-3-amino-piperidin-1-yl)-xanthine (compare WO 2005/085246, example 1 (30)):

1-(2-Cyano-benzyl)-3-methyl-7-(2-butyn-1-yl)-8-((R)-3-amino-piperidin-1-yl)-xanthine (compare WO 2005/085246, example 1 (39)):

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1-[(4-Methyl-quinazolin-2-yl)methyl]-3-methyl-7-(2-butyn-1-yl)-8-[(S)-(2-amino-propyl)-methylamino]-xanthine (compare WO 2006/029769, example 2 (4)):

1-[(3-Cyano-pyridin-2-yl)methyl]-3-methyl-7-(2-butyn-1-yl)-8-((R)-3-amino-piperidin-1-yl)-xanthine (compare WO 2005/085246, example 1 (52)):

1-[(4-Methyl-pyrimidin-2-yl)methyl]-3-methyl-7-(2-butyn-1-yl)-8-((R)-3-amino-piperidin-1-yl)-xanthine (compare WO 2005/085246, example 1 (81)):

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

1-[(4,6-Dimethyl-pyrimidin-2-yl)methyl]-3-methyl-7-(2-butyn-1-yl)-8-((R)-3-amino-piperidin-1-yl)-xanthine (compare WO 2005/085246, example 1 (82)):

1-[(Quinoxalin-6-yl)methyl]-3-methyl-7-(2-butyn-1-yl)-8-((R)-3-amino-piperidin-1-yl)-xanthine (compare WO 2005/085246, example 1 (83)):

A more preferred DPP-4 inhibitor among the abovementioned DPP-4 inhibitors of embodiment A of this invention is 1-[(4-methyl-quinazolin-2-yl)methyl]-3-methyl-7-(2-butyn-1-yl)-8-(3-(R)-amino-piperidin-1-yl)-xanthine, particularly the free base thereof (which is also known as linagliptin or BI 1356).

As further DPP-4 inhibitors the following compounds can be mentioned:

Sitagliptin (MK-0431) having the structural formula A below is (3R)-3-amino-1-[3-(trifluoromethyl)-5,6,7,8-tetrahydro-5H-[1,2,4]triazolo[4,3-a]pyrazin-7-yl]-4-(2, 4,5-trifluorophenyl)butan-1-one, also named (2R)-4-oxo-4-[3-(trifluoromethyl)-5,6-dihydro[1,2,4]triazolo [4,3-a]pyrazin-7(8H)-yl]-1-(2,4,5-trifluorophenyl) butan-2-amine,

In one embodiment, sitagliptin is in the form of its dihydrogenphosphate salt, i.e. sitagliptin phosphate. In a further embodiment, sitagliptin phosphate is in the form of a crystalline anhydrate or monohydrate. A class of this embodiment refers to sitagliptin phosphate monohydrate. Sitagliptin free base and pharmaceutically acceptable salts thereof are disclosed in U.S. Pat. No. 6,699,871 and in Example 7 of WO 03/004498. Crystalline sitagliptin phosphate monohydrate is disclosed in WO 2005/003135 and in WO 2007/050485.

For details, e.g. on a process to manufacture, to formulate or to use this compound or a salt thereof, reference is thus made to these documents.

A tablet formulation for sitagliptin is commercially available under the trade name Januvia®. A tablet formulation for sitagliptin/metformin combination is commercially available under the trade name Janumet®.

Vildagliptin (LAF-237) having the structural formula B below is (25)-{[(3-hydroxyadamantan-1-yl)amino] acetyl}pyrrolidine-2-carbonitrile, also named (S)-1-[(3-hydroxy-1-adamantyl)amino]acetyl-2-cyano-pyrrolidine,

(C)

Vildagliptin is specifically disclosed in U.S. Pat. No. 6,166,063 and in Example 1 of WO 00/34241. Specific salts of vildagliptin are disclosed in WO 2007/019255. A crystalline form of vildagliptin as well as a vildagliptin tablet formulation are disclosed in WO 2006/078593. Vildagliptin can be formulated as described in WO 00/34241 or in WO 2005/067976. A modified release vildagliptin formulation is described in WO 2006/135723.

For details, e.g. on a process to manufacture, to formulate or to use this compound or a salt thereof, reference is thus 10 made to these documents.

A tablet formulation for vildagliptin is commercially available under the trade name Galvus®. A tablet formulation for vildagliptin/metformin combination is commercially available under the trade name Eucreas®.

Saxagliptin (BMS-477118) having the structural formula C below is (1S,3S,5S)-2-{(2S)-2-amino-2-(3-hydroxy-adamantan-1-yl)acetyl}-2-azabicyclo[3.1.0]hexane-3-carbonitrile, also named (S)-3-hydroxyadamantylgly-cine-L-cis-4,5-methanoprolinenitrile,

Saxagliptin is specifically disclosed in U.S. Pat. No. 6,395,767 and in Example 60 of WO 01/68603.

In one embodiment, saxagliptin is in the form of its HCl ³⁵ salt or its mono-benzoate salt as disclosed in WO 2004/052850. In a further embodiment, saxagliptin is in the form of the free base. In a yet further embodiment, saxagliptin is in the form of the monohydrate of the free base as disclosed in WO 2004/052850. Crystalline forms of the HCl salt and of the free base of saxagliptin are disclosed in WO 2008/131149. A process for preparing saxagliptin is also disclosed in WO 2005/106011 and WO 2005/115982. Saxagliptin can be formulated in a tablet as described in WO 2005/117841.

For details, e.g. on a process to manufacture, to formulate ⁴⁵ or to use this compound or a salt thereof, reference is thus made to these documents.

Alogliptin (SYR-322) having the structural formula E below is 2-({6-[(3R)-3-aminopiperidin-1-yl]-3-methyl-2,4-dioxo-3,4-dihydro-2H-pyrimidin-1-yl}methyl) benzonitrile

Alogliptin is specifically disclosed in US 2005/261271, EP 1586571 and in WO 2005/095381. In one embodiment, alogliptin is in the form of its benzoate salt, its hydrochloride salt or its tosylate salt each as disclosed in WO 2007/035629. A class of this embodiment refers to alogliptin benzoate. Polymorphs of alogliptin benzoate are disclosed in WO 2007/035372. A process for preparing alogliptin is disclosed in WO 2007/112368 and, specifically, in WO 2007/035629. Alogliptin (namely its benzoate salt) can be formulated in a tablet and administered as described in WO 2007/033266. A solid preparation of alogliptin/pioglitazone and its preparation and use is described in WO 2008/093882. A solid preparation of alogliptin/metformin and its preparation and use is described in WO 2009/011451. For details, e.g. on a process to manufacture, to formulate or to use this compound or a salt thereof, reference is thus made to these documents.

(2S)-1-{[2-(5-Methyl-2-phenyl-oxazol-4-yl)-ethyl-amino]-acetyl}-pyrrolidine-2-carbonitrile or a pharmaceutically acceptable salt thereof, preferably the mesylate. or

(2S)-1-{[1,1,-Dimethyl-3-(4-pyridin-3-yl-imidazol-1-yl)-propylamino]-acetyl}-pyrrolidine-2-carbonitrile or a pharmaceutically acceptable salt thereof:

These compounds and methods for their preparation are disclosed in WO 03/037327. The mesylate salt of the former compound as well as crystalline polymorphs thereof are disclosed in WO 2006/100181. The fumarate salt of the latter compound as well as crystalline polymorphs thereof are disclosed in WO 2007/071576. These compounds can be formulated in a pharmaceutical composition as described in WO 2007/017423. For details, e.g. on a process to manufacture, to formulate or to use these compounds or salts thereof, reference is thus made to these documents.

(S)-1-((2S,3S,11bS)-2-Amino-9,10-dimethoxy-1,3,4,6,7, 11b-hexahydro-2H-pyrido[2,1-a]isoquinolin-3-yl)-4-fluoromethyl-pyrrolidin-2-one (also named carmegliptin) or a pharmaceutically acceptable salt thereof:

This compound and methods for its preparation are disclosed in WO 2005/000848. A process for preparing this compound (specifically its dihydrochloride salt) is also disclosed in WO 2008/031749, WO 2008/031750 and WO 2008/055814. This compound can be formulated in a pharmaceutical composition as described in WO 2007/017423.

For details, e.g. on a process to manufacture, to formulate 60 or to use this compound or a salt thereof, reference is thus made to these documents.

(3,3-Difluoropyrrolidin-1-yl)-((2S,4S)-4-(4-(pyrimidin-2-yl)piperazin-1-yl)pyrrolidin-2-yl)methanone (also named gosogliptin) or a pharmaceutically acceptable salt thereof:

This compound and methods for its preparation are disclosed in WO 2005/116014 and U.S. Pat. No. 7,291,618.

15

For details, e.g. on a process to manufacture, to formulate or to use this compound or a salt thereof, reference is thus made to these documents.

(1((3S,4S)-4-amino-1-(4-(3,3-difluoropyrrolidin-1-yl)-1, 3,5-triazin-2-yl)pyrrolidin-3-yl)-5,5-difluoropiperidin-2-one or a pharmaceutically acceptable salt thereof:

This compound and methods for its preparation are disclosed in WO 2007/148185 and US 20070299076. For details, e.g. on a process to manufacture, to formulate or to 20 use this compound or a salt thereof, reference is thus made to these documents.

(2S,4S)-1-{2-[(3S,1R)-3-(1H-1,2,4-Triazol-1-ylmethyl) cyclopentylamino]-acetyl}-4-fluoropyrrolidine-2-carcally acceptable salt thereof:

This compound and methods for its preparation are disclosed in WO 2006/040625 and WO 2008/001195. Specifically claimed salts include the methanesulfonate and p-toluenesulfonate. For details, e.g. on a process to manufacture, to formulate or to use this compound or a salt thereof, reference is thus made to these documents.

(R)-2-[6-(3-Amino-piperidin-1-yl)-3-methyl-2,4-dioxo-3,4-dihydro-2H-pyrimidin-1-ylmethyl]-4-fluoro-benzonitrile or a pharmaceutically acceptable salt thereof:

This compound and methods for its preparation and use are disclosed in WO 2005/095381, US 2007060530, WO 2007/033350, WO 2007/035629, WO 2007/074884, WO 2007/112368, WO 2008/033851, WO 2008/114800 and WO 2008/114807. Specifically claimed salts include the succi- 65 nate (WO 2008/067465), benzoate, benzenesulfonate, p-toluenesulfonate, (R)-mandelate and hydrochloride. For

details, e.g. on a process to manufacture, to formulate or to use this compound or a salt thereof, reference is thus made to these documents.

 $5-\{(S)-2-[2-((S)-2-Cyano-pyrrolidin-1-yl)-2-oxo-ethyl$ amino]-propyl}-5-(1H-tetrazol-5-yl)-10,11-dihydro-5H-dibenzo[a,d]cycloheptene-2,8-dicarboxylic bis-dimethylamide or a pharmaceutically acceptable salt thereof:

This compound and methods for its preparation are disbonitrile (also named melogliptin) or a pharmaceuti- 25 closed in WO 2006/116157 and US 2006/270701. For details, e.g. on a process to manufacture, to formulate or to use this compound or a salt thereof, reference is thus made to these documents.

> $3-\{(2S,4S)-4-[4-(3-Methyl-1-phenyl-1H-pyrazol-5-yl)\}$ piperazin-1-yl]pyrrolidin-2-ylcarbonyl}thiazolidine (also named teneligliptin) or a pharmaceutically acceptable salt thereof:

This compound and methods for its preparation are disclosed in WO 02/14271. Specific salts are disclosed in WO 2006/088129 and WO 2006/118127 (including hydrochloride, hydrobromide, inter alia). Combination therapy using this compound is described in WO 2006/129785. For details, e.g. on a process to manufacture, to formulate or to use this compound or a salt thereof, reference is thus made to these documents.

[(2R)-1-{[(3R)-pyrrolidin-3-ylamino]acetyl}pyrrolidin-2-yl]boronic acid (also named dutogliptin) or a pharmaceutically acceptable salt thereof:

This compound and methods for its preparation are dis-45 closed in WO 2005/047297, WO 2008/109681 and WO 2009/009751. Specific salts are disclosed in WO 2008/ 027273 (including citrate, tartrate). A formulation of this compound is described in WO 2008/144730. A formulation of dutogliptin (as its tartrate salt) with metformin is 50 described in WO 2009/091663. For details, e.g. on a process to manufacture, to formulate or to use this compound or a salt thereof, reference is thus made to these documents.

(2S,4S)-1-[2-[(4-ethoxycarbonylbicyclo[2.2.2]oct-1-yl) amino]acetyl]-4-fluoropyrrolidine-2-carbonitrile or a pharmaceutically acceptable salt thereof:

This compound and methods for its preparation are disclosed in WO 2005/075421, US 2008/146818 and WO 2008/114857. For details, e.g. on a process to manufacture, to formulate or to use this compound or a salt thereof, 60 reference is thus made to these documents.

2-({6-[(3R)-3-amino-3-methylpiperidin-1-yl]-1,3-dimethyl-2,4-dioxo-1,2,3,4-tetrahydro-5H-pyrrolo[3,2-d] pyrimidin-5-yl}methyl)-4-fluorobenzonitrile or a pharmaceutically acceptable salt thereof, or 6-[(3R)-3amino-piperidin-1-yl]-5-(2-chloro-5-fluoro-benzyl)-1, 3-dimethyl-1,5-dihydro-pyrrolo[3,2-d]pyrimidine-2,4dione or a pharmaceutically acceptable salt thereof:

triple combination therapy using the DPP-4 inhibitor, the second and the third antidiabetic agent.

30

These compounds and methods for their preparation are disclosed in WO 2009/084497 and WO 2006/068163, respectively. Combination therapy using the latter of these two compounds is described in WO 2009/128360. For details, e.g. on a process to manufacture, to formulate or to 5 use these compounds or salts thereof, reference is thus made to these documents.

(S)-2-methylpyrazolo[1,5-a]primidine-6-carboxylic acid {2-[(2-cyanopyrrolidin-1-yl)-2-oxoethylamino]-2-methylpropyl}amide (also named anagliptin) or a pharmaceutically acceptable salt:

This compound and methods for its preparation are disclosed in WO 2004/067509. Combination therapy using this compound is described in WO 2009/139362. For details, e.g. on a process to manufacture, to formulate or to use this 15 compound or a salt thereof, reference is thus made to these documents.

Preferably the DPP-4 inhibitor is selected from the group G2 consisting of linagliptin, sitagliptin, vildagliptin, alogliptin, saxagliptin, carmegliptin, gosogliptin, teneligliptin, 20 melogliptin and dutogliptin, or a pharmaceutically acceptable salt of one of the hereinmentioned DPP-4 inhibitors, or a prodrug thereof.

More preferably the DPP-4 inhibitor is selected from the group G2 consisting of linagliptin, sitagliptin, vildagliptin, 25 alogliptin, saxagliptin, teneligliptin and dutogliptin, or a pharmaceutically acceptable salt of one of the hereinmentioned DPP-4 inhibitors, or a prodrug thereof.

A particularly preferred DPP-4 inhibitor within the present invention is linagliptin. The term "linagliptin" as 30 employed herein refers to linagliptin and pharmaceutically acceptable salts thereof, including hydrates and solvates thereof, and crystalline forms thereof. Crystalline forms are described in WO 2007/128721. Methods for the manufacture of linagliptin are described in the patent applications 35 WO 2004/018468 and WO 2006/048427 for example. Linagliptin is distinguished from structurally comparable DPP-4 inhibitors, as it combines exceptional potency and a longlasting effect with favourable pharmacological properties, receptor selectivity and a favourable side-effect profile or 40 bring about unexpected therapeutic advantages or improvements in monotherapy and/or when used in combination with a second and, optionally, a third antidiabetic agent according to this invention.

For avoidance of any doubt, the disclosure of each of the 45 foregoing documents cited above in connection with the specified DPP-4 inhibitors is specifically incorporated herein by reference in its entirety.

In one aspect of the present invention, the pharmaceutical compositions, methods and uses according to this invention 50 relate to those compositions which comprise the DPP-4 inhibitor as sole active ingredient (i.e. the second and third antidiabetic agent are both absent) and/or, respectively, to monotherapy using the DPP-4 inhibitor alone.

In another aspect of the present invention, the pharmaceutical compositions, combinations, methods and uses according to this invention relate to those compositions or combinations which comprise the DPP-4 inhibitor and the second antidiabetic agent as sole active ingredients (i.e. the third antidiabetic agent is absent) and/or, respectively, to 60 dual combination therapy using the DPP-4 inhibitor and the second antidiabetic agent.

In another aspect of the present invention, the pharmaceutical compositions, combinations, methods and uses according to this invention relate to those compositions or 65 combinations which comprise the DPP-4 inhibitor, the second and the third antidiabetic agent and/or, respectively, to

Further, a DPP-4 inhibitor according to this invention may be further characterized in that said DPP-4 inhibitor does not significantly impair glomerular and/or tubular function of a type 2 diabetes patient with chronic renal insufficiency (e.g. mild, moderate or severe renal impairment or end stage renal disease), and/or

said DPP-4 inhibitor does not require to be dose-adjusted in a type 2 diabetes patient with impaired renal function (e.g. mild, moderate or severe renal impairment or end stage renal disease).

The second antidiabetic agent and, if present, the third antidiabetic agent is selected from the group G3 consisting of biguanides, thiazolidindiones, sulfonylureas, glinides, inhibitors of alpha-glucosidase, GLP-1 or GLP-1 analogues, and insulin or insulin analogues, or a pharmaceutically acceptable salt thereof. In the following preferred embodiments regarding the second and/or the third antidiabetic agent are described.

The group G3 comprises biguanides. Examples of biguanides are metformin, phenformin and buformin. A preferred biguanide is metformin. A DPP-4 inhibitor in combination with a biguanide, in particular metformin, can provide more efficacious glycemic control and/or may act together with the biguanide, for example to reduce weight, that has e.g. overall beneficial effects on the metabolic syndrome which is commonly associated with type 2 diabetes mellitus.

The term "metformin" as employed herein refers to metformin or a pharmaceutically acceptable salt thereof such as the hydrochloride salt, the metformin (2:1) fumarate salt, and the metformin (2:1) succinate salt, the hydrobromide salt, the p-chlorophenoxy acetate or the embonate, and other known metformin salts of mono and dibasic carboxylic acids. It is preferred that the metformin employed herein is the metformin hydrochloride salt.

The group G3 comprises thiazolidindiones. Examples of thiazolidindiones (TZD) are pioglitazone and rosiglitazone. TZD therapy is associated with weight gain and fat redistribution. In addition, TZD cause fluid retention and are not indicated in patients with congestive heart failure. Long term treatment with TZD are further associated with an increased risk of bone fractures. A DPP-4 inhibitor in combination with a thiazolidindione, in particular pioglitazone, can provide more efficacious glycemic control and/or can minimize side effects of the treatment with TZD.

The term "pioglitazone" as employed herein refers to pioglitazone, including its enantiomers, mixtures thereof and its racemate, or a pharmaceutically acceptable salt thereof such as the hydrochloride salt.

The term "rosiglitazone" as employed herein refers to rosiglitazone, including its enantiomers, mixtures thereof and its racemate, or a pharmaceutically acceptable salt thereof such as the maleate salt.

The group G3 comprises sulfonylureas. Examples of sulfonylureas are glibenclamide, tolbutamide, glimepiride, glipizide, gliquidone, glibornuride, glyburide, glisoxepide and gliclazide. Preferred sulfonylureas are tolbutamide, gliquidone, glibenclamide and glimepiride, in particular glibenclamide and glimepiride. As the efficacy of sulfonylureas wears off over the course of treatment, a combination of a DPP-4 inhibitor with a sulfonylurea may offer additional benefit to the patient in terms of better glycemic control. Also, treatment with sulfonylureas is normally associated with gradual weight gain over the course of treatment and a DPP-4 inhibitor may minimize this side effect of the treatment with an sulfonylurea and/or improve the metabolic

32 TABLE 1

syndrome. Also, a DPP-4 inhibitor in combination with a
sulfonylurea may minimize hypoglycemia which is another
undesirable side effect of sulfonylureas. This combination
may also allow a reduction in the dose of sulfonylureas,
which may also translate into less hypoglycemia.
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Each term of the group "glibenclamide", "glimepiride", "gliquidone", "glibornuride", "gliclazide", "glisoxepide", "tolbutamide" and "glipizide" as employed herein refers to the respective active drug or a pharmaceutically acceptable salt thereof

The group G3 comprises glinides. Examples of glinides are nateglinide, repaglinide and mitiglinide. As their efficacy wears off over the course of treatment, a combination of a DPP-4 inhibitor with a meglitinide may offer additional benefit to the patient in terms of better glycemic control. Also, treatment with meglitinides is normally associated with gradual weight gain over the course of treatment and a DPP-4 inhibitor may minimize this side effect of the treatment with an meglitinide and/or improve the metabolic 20 syndrome. Also, a DPP-4 inhibitor in combination with a meglitinide may minimize hypoglycemia which is another undesirable side effect of meglitinides. This combination may also allow a reduction in the dose of meglitinides, which may also translate into less hypoglycemia.

The term "nateglinide" as employed herein refers to nateglinide, including its enantiomers, mixtures thereof and its racemate, or a pharmaceutically acceptable salts and esters thereof.

The term "repaglinide" as employed herein refers to ³⁰ repaglinide, including its enantiomers, mixtures thereof and its racemate, or a pharmaceutically acceptable salts and esters thereof.

The group G3 comprises inhibitors of alpha-glucosidase. Examples of inhibitors of alpha-glucosidase are acarbose, voglibose and miglitol. Additional benefits from the combination of a DPP-4 inhibitor and an alpha-glucosidase inhibitor may relate to more efficacious glycemic control, e.g. at lower doses of the individual drugs, and/or reducement of undesirable gastrointestinal side effects of alpha-glucosidase inhibitors.

Each term of the group "acarbose", "voglibose" and "miglitol" as employed herein refers to the respective active drug or a pharmaceutically acceptable salt thereof.

The group G3 comprises inhibitors of GLP-1 analogues. Examples of GLP-1 analogues are exenatide, liraglutide, taspoglutide, semaglutide, albiglutide, and lixisenatide. The combination of a DPP-4 inhibitor and a GLP-1 analogue may achieve a superior glycemic control, e.g. at lower doses of the individual drugs. In addition, e.g. the body weight reducing capability of the GLP-1 analogue may be positively act together with the properties of the DPP-4 inhibitor. On the other hand, a reduction of side effects (e.g. nausea, gastrointestinal side effects like vomiting) may be obtained, e.g. when a reduced dose of the GLP-1 analogue is applied in the combination with a DPP-4 inhibitor.

Each term of the group "exenatide", "liraglutide", "taspoglutide", "semaglutide", "albiglutide", and "lixisenatide" as $_{60}$ employed herein refers to the respective active drug or a pharmaceutically acceptable salt thereof.

In an embodiment (embodiment E1) the pharmaceutical compositions, combinations, methods and uses according to this invention relate to those combinations wherein the 65 DPP-4 inhibitor and the second antidiabetic agent are preferably selected according to the entries in the Table 1.

DPP-4 Inhibitor	Second Antidiabetic Agent
selected from embodiment B	selected from the group G3
selected from embodiment B	Metformin
selected from embodiment B	Pioglitazone
selected from embodiment B selected from embodiment B	Rosiglitazone Glibenclamide
selected from embodiment B	Glimepiride
selected from embodiment B	Gliquidone
selected from embodiment B	Nateglinide
selected from embodiment B	Repaglinide
selected from embodiment B selected from embodiment B	Acarbose
selected from embodiment B	Voglibose Miglitol
selected from embodiment B	Exenatide
selected from embodiment B	Liraglutide
selected from embodiment B	Taspoglutide
selected from embodiment B selected from embodiment B	Semaglutide
selected from embodiment B	Albiglutide Lixisenatide
Linagliptin	selected from the group G3
Linagliptin	Metformin
Linagliptin	Pioglitazone
Linagliptin	Rosiglitazone
Linagliptin Linagliptin	Glibenclamide Glimepiride
Linagliptin	Gliquidone
Linagliptin	Nateglinide
Linagliptin	Repaglinide
Linagliptin	Acarbose
Linagliptin Linagliptin	Voglibose
Linagliptin	Miglitol Exenatide
Linagliptin	Liraglutide
Linagliptin	Taspoglutide
Linagliptin	Semaglutide
Linagliptin	Albiglutide Lixisenatide
Linagliptin Sitagliptin	selected from the group G3
Sitagliptin	Metformin
Sitagliptin	Pioglitazone
Sitagliptin	Rosiglitazone
Sitagliptin Sitagliptin	Glibenclamide Glimepiride
Sitagliptin	Gliquidone
Sitagliptin	Nateglinide
Sitagliptin	Repaglinide
Sitagliptin Sitagliptin	Acarbose Voglibose
Sitagliptin	Miglitol
Sitagliptin	Exenatide
Sitagliptin	Liraglutide
Sitagliptin	Taspoglutide Semaglutide
Sitagliptin Sitagliptin	Albiglutide
Sitagliptin	Lixisenatide
Vildagliptin	selected from the group G3
Vildagliptin	Metformin
Vildagliptin	Pioglitazone Rosiglitazone
Vildagliptin Vildagliptin	Glibenclamide
Vildagliptin	Glimepiride
Vildagliptin	Gliquidone
Vildagliptin	Nateglinide
Vildagliptin	Repaglinide
Vildagliptin Vildagliptin	Acarbose Voglibose
Vildagliptin	Miglitol
Vildagliptin	Exenatide
Vildagliptin	Liraglutide
Vildagliptin	Taspoglutide Samaglutida
Vildagliptin Vildagliptin	Semaglutide Albiglutide
Vildagliptin	Lixisenatide
Alogliptin	selected from the group G3
Alogliptin	Metformin
Alogliptin	Pioglitazone Pogialitazone
Alogliptin Alogliptin	Rosiglitazone Glibenclamide
Alogliptin	Glimepiride
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34
TABLE 1-continued

TAB	LE 1-continued	_	TAI	BLE 1-continued
DPP-4 Inhibitor	Second Antidiabetic Agent		DPP-4 Inhibitor	Second Antidiabetic Agent
Alogliptin	Gliquidone	_	Dutogliptin	Exenatide
Alogliptin	Nateglinide	5	Dutogliptin	Liraglutide
Alogliptin	Repaglinide		Dutogliptin	Taspoglutide
Alogliptin	Acarbose		Dutogliptin	Semaglutide
Alogliptin	Voglibose		Dutogliptin	Albiglutide
Alogliptin	Miglitol		Dutogliptin	Lixisenatide
Alogliptin	Exenatide		Gosogliptin	selected from the group G3
Alogliptin	Liraglutide	10	Gosogliptin	Metformin
Alogliptin	Taspoglutide		Gosogliptin	Pioglitazone
Alogliptin	Semaglutide		Gosogliptin	Rosiglitazone
Alogliptin	Albiglutide		Gosogliptin	Glibenclamide
Alogliptin	Lixisenatide		Gosogliptin	Glimepiride
Saxagliptin	selected from the group G3		Gosogliptin	Gliquidone Nateglinide
Saxagliptin	Metformin	15	Gosogliptin	
Saxagliptin	Pioglitazone		Gosogliptin	Repaglinide Acarbose
Saxagliptin Saxagliptin	Rosiglitazone		Gosogliptin Gosogliptin	Voglibose
Saxaglipun Saxagliptin	Glibenclamide Glimepiride			Vogitoose Miglitol
			Gosogliptin	
Saxagliptin	Gliquidone		Gosogliptin	Exenatide
Saxagliptin	Nateglinide	20	Gosogliptin	Liraglutide
Saxagliptin	Repaglinide		Gosogliptin	Taspoglutide
Saxagliptin	Acarbose		Gosogliptin	Semaglutide
Saxagliptin	Voglibose		Gosogliptin	Albiglutide
Saxagliptin	Miglitol		Gosogliptin	Lixisenatide
Saxagliptin	Exenatide		Teneligliptin	selected from the group G3
Saxagliptin	Liraglutide	25	Teneligliptin	Metformin
Saxagliptin	Taspoglutide	25	Teneligliptin	Pioglitazone
Saxagliptin	Semaglutide		Teneligliptin	Rosiglitazone
Saxagliptin	Albiglutide		Teneligliptin	Glibenclamide
Saxagliptin	Lixisenatide		Teneligliptin	Glimepiride
Carmegliptin	selected from the group G3		Teneligliptin	Gliquidone
Carmegliptin	Metformin		Teneligliptin	Nateglinide
Carmegliptin	Pioglitazone	30	Teneligliptin	Repaglinide
Carmegliptin	Rosiglitazone		Teneligliptin	Acarbose
Carmegliptin	Glibenclamide		Teneligliptin	Voglibose
Carmegliptin	Glimepiride		Teneligliptin	Miglitol
Carmegliptin	Gliquidone		Teneligliptin	Exenatide
Carmegliptin	Nateglinide		Teneligliptin	Liraglutide
Carmegliptin	Repaglinide	35	Teneligliptin	Taspoglutide
Carmegliptin	Acarbose	33	Teneligliptin	Semaglutide
Carmegliptin	Voglibose		Teneligliptin	Albiglutide
Carmegliptin	Miglitol		Teneligliptin	Lixisenatide
Carmegliptin	Exenatide			
Carmegliptin	Liraglutide			
Carmegliptin	Taspoglutide		In a particular embo	odiment (embodiment E2) the phar
Carmegliptin	Semaglutide	40		ns, combinations, methods and use
Carmegliptin	Albiglutide			
Carmegliptin	Lixisenatide		according to this inve	ention relate to those combination
Melogliptin	selected from the group G3		wherein the DPP-4 in	hibitor is linagliptin. According to
Melogliptin	Metformin			
Melogliptin	Pioglitazone			cond antidiabetic agent is preferably
Melogliptin	Rosiglitazone	45	selected according to the	he entries in the Table 2.
Melogliptin	Glibenclamide		2	
Melogliptin	Glimepiride			TABLE 2
Melogliptin	Gliquidone			TABLE 2
Melogliptin Melogliptin	Nateglinide			
			Embodiment	Second Antidiabetic Agent
Melogliptin Melogliptin	Repaglinide			1 / 10 / 2
Melogliptin Melogliptin	Acarbose Vanlihaga	50	E2.1	selected from the group G3
Melogliptin	Voglibose		E2.2	Metformin
Melogliptin	Miglitol		E2.3	Pioglitazone
Melogliptin	Exenatide		E2.4	Rosiglitazone
Melogliptin	Liraglutide		E2.5	Glibenclamide
Melogliptin	Taspoglutide		E2.6	Glimepiride
Melogliptin	Semaglutide	55	E2.7	Gliquidone
Melogliptin	Albiglutide		E2.8	Nateglinide
Melogliptin	Lixisenatide		E2.9	Repaglinide
Dutogliptin	selected from the group G3		E2.10	Acarbose
Dutogliptin	Metformin		E2.11	Voglibose
Dutogliptin	Pioglitazone		E2.12	Miglitol
Dutogliptin	Rosiglitazone	CO	E2.13	Exenatide
Dutogliptin	Glibenclamide	60	E2.14	Liraglutide
Dutogliptin	Glimepiride		E2.15	Taspoglutide
OI	Gliquidone		E2.16	Semaglutide
	Gilquidolle			
Dutogliptin	Nateglinide		E2.17	Albiglutide
Dutogliptin Dutogliptin Dutogliptin			E2.17 E2.18	Albiglutide Lixisenatide
Dutogliptin Dutogliptin	Nateglinide		E2.18	Lixisenatide
Dutogliptin Dutogliptin Dutogliptin	Nateglinide Repaglinide	65		

The combination of a DPP-4 inhibitor and a second and, optionally, a third antidiabetic agent according to this invention can be found to improve the glycemic control, in particular in patients as described herein, compared with a monotherapy using either a DPP-4 inhibitor or the second or 5 third antidiabetic agent alone, for example with a monotherapy of metformin, or with a dual therapy using the second and third antidiabetic agent. Further, the triple combination of a DPP-4 inhibitor and a second and a third antidiabetic agent according to this invention can be found 10 to improve the glycemic control, in particular in patients as described herein, compared with a combination therapy using a DPP-4 inhibitor and either the second or third antidiabetic agent, or using the second and the third antidiabetic agent. The improved glycemic control is determined 15 as an increased lowering of blood glucose and an increased reduction of HbA1c. With monotherapy in a patient, in particular in patients as described herein, the glycemic control may not be further improved significantly by an administration of the drug above a certain highest dose. In 20 addition, a long term treatment using a highest dose may be unwanted in view of potential side effects. Therefore, a satisfying glycemic control may not be achievable in all patients via a monotherapy using either the DPP-4 inhibitor or the second or the third antidiabetic agent alone. In the case 25 that monotherapy do not yield in full glycemic control, dual therapy may become necessary. Even with combination therapy using two agents selected from the DPP-4 inhibitors and second and third antidiabetic agents may not yield in a full glycemic control in all patients and/or over a long time. 30 In the case that dual therapy do not yield in full glycemic control, triple therapy may become necessary. In such patients with inadequate glycemic control a progression of the diabetes mellitus may continue and complications associated with diabetes mellitus may occur, such as macrovas- 35 cular complications. The pharmaceutical composition or combination as well as the methods according to the present invention allow a reduction of the HbA1c value to a desired target range, for example <7% and preferably <6.5%, for a higher number of patients and for a longer time of thera- 40 peutic treatment, e.g. in the case of dual or triple combination therapy compared with a monotherapy using one of or, respectively, a dual therapy using two of the combination partners.

In addition, the combination of a DPP-4 inhibitor and the 45 second and, optionally, the third therapeutic agent according to this invention can be found to allow a reduction in the dose of either the DPP-4 inhibitor or the second or third antidiabetic agent or even of two or three of the active ingredients. A dose reduction is beneficial for patients which 50 otherwise would potentially suffer from side effects in a therapy using a higher dose of one or more of the active ingredients, in particular with regard to side effect caused by the second and/or third antidiabetic agent. Therefore, the pharmaceutical combination as well as the methods according to the present invention, may show less side effects, thereby making the therapy more tolerable and improving the patients compliance with the treatment.

A DPP-4 inhibitor according to the present invention is able—via the increases in active GLP-1 levels—to reduce 60 the glucagon secretion in a patient. This will therefore limit the hepatic glucose production. Furthermore, the elevated active GLP-1 levels produced by the DPP-4 inhibitor will have beneficial effects on beta-cell regeneration and neogenesis. All these features of DPP-4 inhibitors may render a 65 pharmaceutical composition or combination or method of this invention quite useful and therapeutically relevant.

36

When this invention refers to patients requiring treatment or prevention, it relates primarily to treatment and prevention in humans, but the pharmaceutical composition may also be used accordingly in veterinary medicine in mammals. In the scope of this invention adult patients are preferably humans of the age of 18 years or older. Also in the scope of this invention, patients are adolescent humans, i.e. humans of age 10 to less than 18 years, preferably of age 13 to less than 18 years.

In one embodiment, patients in need of treatment or prevention as described herein can be identified by determining whether they have variation(s) (e.g. polymorphisms) in one or more genes associated with metabolic diseases and/or whether they have variation(s) (e.g. polymorphisms) in one or more of the genes selected from TCF7L2, KCNJ11, PPARG and GLP1R, in particular whether they are of TCF7L2 risk genotype as described herein.

In another embodiment, patients in need of treatment or prevention as described herein can be identified by determining whether they are of respective wild-type genotype, in particular whether they are of TCF7L2 wild genotype as described herein.

A particular sub-population of the patients in need of treatment or prevention as described herein, refers to those patients who have one or more single nucleotide polymorphisms (SNPs) in the gene coding for TCF7L2, especially a SNP selected from rs7903146, rs12255372 and rs10885406, especially rs7903146, in more particular, those patients who carry at least one T allele of SNP rs7903146 of TCF7L2, i.e. the CT genotype or TT genotype.

Another particular sub-population of the patients in need of treatment or prevention as described herein, refers to those patients who carry TCF7L2 rs7903146 CC wild genotype.

Thus, in an aspect of this invention, a treatment or prophylaxis according to this invention is suitable in those patients in need of such treatment or prophylaxis who are diagnosed of having variation(s) (e.g. polymorphisms) in one or more genes associated with metabolic diseases and/or variation(s) (e.g. SNPs) in one or more of the genes selected from TCF7L2, KCNJ11, PPARG and GLP1R, in particular of TCF7L2 risk genotype as described herein.

In another aspect of this invention, a treatment or prophylaxis according to this invention is particular suitable in those patients in need of such treatment or prophylaxis who are diagnosed of having TCF7L2 wild genotype as described herein.

In an sub-aspect of this invention, a treatment or prophylaxis according to this invention is suitable in those patients in need of such treatment or prophylaxis who are diagnosed of having one or more single nucleotide polymorphisms (SNPs) in the gene coding for TCF7L2, e.g. at least one SNP selected from rs7903146, rs12255372 and rs10885406, for example rs7903146, in particular, carrying at least one T allele of rs7903146, (i.e. of CT or TT genotype), among them, in more particular, those carrying one T allele of rs7903146 (i.e. of CT risk genotype) or, in less particular, those carrying two T alleles of rs7903146 (i.e. of TT high risk genotype).

In another sub-aspect of this invention, a treatment or prophylaxis according to this invention is particular favorable in those patients in need of such treatment or prophylaxis who are diagnosed of carrying wild-type two C alleles of rs7903146 in TCF7L2 (i.e. of CC genotype).

In an embodiment of this invention, a treatment or prophylaxis according to this invention is suitable in those patients in need of such treatment or prophylaxis who are

diagnosed of one or more of the conditions selected from the group consisting of overweight and obesity, in particular class I obesity, class II obesity, class III obesity, visceral obesity and abdominal obesity. In addition a treatment or prophylaxis according to this invention is advantageously 5 suitable in those patients in which a weight increase is contraindicated. Any weight increasing effect in the therapy, for example due to the administration of the second and/or third antidiabetic agent, may be attenuated or even avoided thereby.

In a further embodiment of this invention, the pharmaceutical composition or combination of this invention exhibits a very good efficacy with regard to glycemic control, in particular in view of a reduction of fasting plasma glucose, postprandial plasma glucose and/or glycosylated hemoglo- 15 bin (HbA1c). By administering a pharmaceutical composition or combination according to this invention, a reduction of HbA1c equal to or greater than preferably 1.0%, more preferably equal to or greater than 2.0%, even more preferably equal to or greater than 3.0% can be achieved and the 20 reduction is particularly in the range from 1.0% to 3.0%.

Furthermore, the method and/or use according to this invention is applicable in those patients who show one, two or more of the following conditions:

- (a) a fasting blood glucose or serum glucose concentration 25 greater than 110 mg/dL or greater than 100 mg/dL, in particular greater than 125 mg/dL;
- (b) a postprandial plasma glucose equal to or greater than 140 mg/dL;
- (c) an HbA1c value equal to or greater than 6.5%, in 30 particular equal to or greater than 7.0%, especially equal to or greater than 7.5%, even more particularly equal to or greater than 8.0%.

The present invention also discloses the use of the pharmaceutical composition or combination for improving gly- 35 cemic control in patients having type 2 diabetes or showing first signs of pre-diabetes. Thus, the invention also includes diabetes prevention. If therefore a pharmaceutical composition or combination of this invention is used to improve the glycemic control as soon as one of the above-mentioned 40 include, without being limited, patients of Caucasian, North signs of pre-diabetes is present, the onset of manifest type 2 diabetes mellitus can be delayed or prevented.

Furthermore, the pharmaceutical composition or combination of this invention is particularly suitable in the treatment of patients with insulin dependency, i.e. in patients 45 who are treated or otherwise would be treated or need treatment with an insulin or a derivative of insulin or a substitute of insulin or a formulation comprising an insulin or a derivative or substitute thereof. These patients include patients with diabetes type 2 and patients with diabetes type 50

Therefore, according to an embodiment of the present invention, there is provided a method for improving glycemic control and/or for reducing of fasting plasma glucose, of postprandial plasma glucose and/or of glycosylated hemo- 55 globin HbA1c in a patient in need thereof who is diagnosed with impaired glucose tolerance (IGT), impaired fasting blood glucose (IFG) with insulin resistance, with metabolic syndrome and/or with type 2 or type 1 diabetes mellitus characterized in that a DPP-4 inhibitor and, optionally, a 60 second and, optionally, a third antidiabetic agent as defined hereinbefore and hereinafter are administered, for example in combination, to the patient.

According to another embodiment of the present invention, there is provided a method for improving glycemic 65 control in patients, in particular in adult patients, with type 2 diabetes mellitus as an adjunct to diet and exercise.

38

Moreover, in a particular embodiment of this invention, a therapeutic or preventive method and/or use according to this invention is suitable in those patients who have variation(s) (e.g. polymorphisms) in one or more genes associated with metabolic diseases and/or who have variation(s) (e.g. polymorphisms) in one or more of the genes selected from TCF7L2, KCNJ11, PPARG and GLP1R.

In this context, a sub-population of the patients described hereinbefore and hereinafter refers to TCF7L2 risk genotype patients, such as e.g. to those patients who have one or more single nucleotide polymorphisms (SNPs) in the gene coding for TCF7L2, especially at least one SNP selected from rs7903146, rs12255372 and rs10885406, especially rs7903146. In more particular, those patients who carry at least one T allele of SNP rs7903146 of TCF7L2, i.e. the CT genotype or TT genotype, especially who carry two T alleles of SNP rs7903146 of TCF7L2, i.e. the TT genotype, are strongly susceptible to increased TCF7L2 expression in pancreatic beta cells, impaired insulin secretion, incretine effects, enhanced rate of hepatic glucose production and/or diabetes. The T allele of rs7903146 TCF7L2 is associated with impaired insulinotropic action of incretin hormones, reduced 24 h profiles of plasma insulin and glucagon, and increased hepatic glucose production.

Therefore, the present invention also includes the compounds, pharmaceutical compositions or combinations according to this invention for use in the treatment and/or prevention of those diseases, disorders or conditions mentioned herein in those patients who have one or more single nucleotide polymorphisms (SNPs) in the gene coding for TCF7L2, especially at least one SNP selected from rs7903146, rs12255372 and rs10885406, especially rs7903146; in more particular, in those patients who carry at least one T allele of SNP rs7903146 of TCF7L2, i.e. the CT genotype or TT genotype, particularly in those patients who carry one T allele of SNP rs7903146 of TCF7L2, i.e. the CT genotype, or who carry two T alleles of SNP rs7903146 of TCF7L2, i.e. the TT genotype.

TCF7L2 risk genotype patients as described herein European, East Asian, Indian and/or African descent.

The present invention further includes a therapeutic and/ or preventive method or use according to this invention for application in a patient in need thereof, said method or use comprising the step of determining whether the patient has variation(s) (e.g. polymorphisms) in one or more of the genes selected from TCF7L2, KCNJ11, PPARG and GLP1R, particularly whether the patient is of a TCF7L2 risk genotype as described herein.

The determination or diagnosis whether the patient has variation(s) (e.g. polymorphisms) in one or more of the genes selected from TCF7L2, KCNJ11, PPARG and GLP1R, particularly whether the patient is of a TCF7L2 risk genotype as described herein, or whether the patient is of wild genotype, particularly whether the patient is of TCF7L2 wild genotype as described herein, may be used for determining the likelihood (e.g., increased, decreased, or no likelihood) of a favourable therapeutic and/or preventive response of the patient to the treatment with a DPP-4 inhibitor (or with a combination of a DPP-4 inhibitor with the second and/or third antidiabetic agent as defined herein) in a therapeutic and/or preventive method or use as described hereinabove or hereinbelow (e.g. in treating diabetes or in improving glycemic control), and thus for identifying a subject being susceptible to such treatment.

Thus, further on, in another embodiment of this invention, there is provided a method of determining the probability of

likelihood (e.g., increased, decreased, or no likelihood) of a favourable response to the administration of a pharmaceutically acceptable amount of a DPP-4 inhibitor (or of a combination of a DPP-4 inhibitor with the second and/or third antidiabetic agent as described herein) in a subject (particularly diabetes patient), said method comprising the step of determining whether the subject has variation(s) (e.g. polymorphisms) in one or more of the genes selected from TCF7L2, KCNJ11, PPARG and GLP1R, particularly whether the subject is of a TCF7L2 risk genotype as described herein, or determining whether the subject is of TCF7L2 wild genotype, particularly testing whether the subject is of the TCF7L2 rs7903146 CC wild genotype.

According to another particular embodiment this invention, the present invention provides a DPP-4 inhibitor, a pharmaceutical composition or combination according to the present invention for use in a therapeutic or preventive method as described hereinbefore or hereinafter (particularly for treating or preventing type 2 diabetes and/or 20 obesity), said method comprising

(i) identifying a subject being susceptible to the therapeutic or preventive method, said identifying comprising testing whether the subject has variation(s) (e.g. polymorphisms) in one or more of the genes selected from TCF7L2, KCNJ11, 25 PPARG and GLP1R, in particular whether the subject is of any TCF7L2 risk genotype as described herein, in more particular whether he/she has one or more single nucleotide polymorphisms (SNPs) in the gene coding for TCF7L2, especially at least one SNP selected from rs7903146, 30 rs12255372 and rs10885406, especially rs7903146, for example whether the subject carries at least one T allele of SNP rs7903146 of TCF7L2, e.g. whether the subject is of CT genotype (i.e. whether the patient carries one T allele of SNP rs7903146 of TCF7L2) or whether the subject is of TT 35 genotype (i.e. whether the patient carries two T alleles of SNP rs7903146 of TCF7L2), or testing whether the subject is of TCF7L2 wild genotype, in particular whether the subject is of the TCF7L2 rs7903146 CC wild genotype; and thus determining the probability of likelihood of a favour- 40 able response (e.g. favorable change in HbA1c) resulting from the rapeutic or preventive treatment of the subject with the DPP-4 inhibitor, pharmaceutical composition or combi-

and

(ii) administering an effective amount of the DPP-4 inhibitor, pharmaceutical composition or combination to the subject, where said subject is determined to have a high probability of likelihood of a favorable response (e.g. favorable change in HbA1c) resulting from therapeutic or preventive treatment with the DPP-4 inhibitor, pharmaceutical composition or combination.

The present invention further provides a therapeutic and/ or preventive method or use of this invention for application in a patient in need thereof, said method or use comprising 55 the steps of

obtaining and assaying a nucleic acid sample from an individual with type 2 diabetes mellitus,

determining the efficacy and/or, optionally, the probability of the likelihood of a favorable response (e.g. in 60 providing glycemic control, such as favorable change in HbA1c) to a treatment with a DPP-4 inhibitor, preferably linagliptin, or the DPP-4 inhibitor in combination with one or more other active substances (e.g. antidiabetics), comprising detecting either TT or CT or CC allele genotype at rs7903146 of TCF7L2 gene in patient's sample,

40

wherein the presence of the TT, CT or CC genotype is indicative of the efficacy to the treatment, and/or, optionally,

wherein the presence of the TT genotype is indicative of a decreased likelihood of favorable response and/or presence of the CC genotype is indicative of an increased likelihood of favorable response to the treatment, and

administering a therapeutically effective amount of the DPP-4 inhibitor, preferably linagliptin, or the DPP-4 inhibitor in combination with one or more other active substances (e.g. antidiabetics) to the individual.

It can be further found that by using a pharmaceutical composition or combination according to this invention, an improvement of the glycemic control can be achieved even in those patients who have insufficient glycemic control in particular despite treatment with the second or third antidiabetic agent or a combination of the second with the third antidiabetic agent, for example despite maximal tolerated dose of oral monotherapy with metformin, a thiazolidinedione (e.g. pioglitazone) or a sulfonylurea, or a combination of metformin with a thiazolidinedione (e.g. pioglitazone), of metformin with a sulfonylurea, or of a thiazolidinedione (e.g. pioglitazone) with a sulfonylurea.

It can be also found that by using a combination according to this invention, an improvement of the glycemic control can be achieved even in those patients who have insufficient glycemic control in particular despite treatment with a DPP-4 inhibitor or a combination of a DPP-4 inhibitor with the second or third antidiabetic agent, for example despite maximal tolerated dose of oral monotherapy with a DPP-4 inhibitor or a dual combination of a DPP-4 inhibitor with the second or third antidiabetic agent.

A maximal tolerated dose with regard to metformin is for example 2000 mg per day, 1500 mg per day (for example in asian countries) or 850 mg three times a day or any equivalent thereof.

Therefore, the method and/or use according to this invention is applicable in those patients who show one, two or more of the following conditions:

- (a) insufficient glycemic control with diet and exercise alone:
- (b) insufficient glycemic control despite monotherapy with metformin, a thiazolidinedione (e.g. pioglitazone), a sulfonylurea, GLP-1 or GLP-1 analogue, or insulin or insulin analogue, in particular despite oral monotherapy at a maximal tolerated dose of metformin, a thiazolidinedione (e.g. pioglitazone) or a sulfonylurea;
- (c) insufficient glycemic control despite combination therapy with two agents selected from the group consisting of metformin, a thiazolidinedione (e.g. pioglitazone), a sulfonylurea, GLP-1 or GLP-1 analogue, and insulin or insulin analogue, for example despite combination therapy with a dual combination selected from metformin/ pioglitazone, metformin/sulphonylurea, metformin/insulin, sulphonylurea/pioglitazone, sulphonylurea/insulin and pioglitazone/insulin;

The dual or triple combination method and/or use according to this invention is further applicable in those patients who show the following conditions (e) or (f), respectively: (d) insufficient glycemic control despite oral monotherapy with the DPP-4 inhibitor, in particular despite oral monotherapy at a maximal tolerated dose of the DPP-4 inhibitor:

(e) insufficient glycemic control despite (oral) combination therapy with the DPP-4 inhibitor and the second or third antidiabetic agent, in particular despite oral dual therapy at a maximal tolerated dose of at least one of the combination partners.

In an embodiment of this invention, a pharmaceutical composition or combination is suitable in the treatment of patients who are diagnosed having one or more of the following conditions

insulin resistance,

hyperinsulinemia,

pre-diabetes,

type 2 diabetes mellitus, particular having a late stage type 2 diabetes mellitus,

type 1 diabetes mellitus.

Furthermore, a pharmaceutical composition or combination according to this invention is particularly suitable in the treatment of patients who are diagnosed having one or more of the following conditions

- (a) obesity (including class I, II and/or III obesity), visceral obesity and/or abdominal obesity,
- (b) triglyceride blood level ≥150 mg/dL,
- (c) HDL-cholesterol blood level <40 mg/dL in female patients and <50 mg/dL in male patients,
- (d) a systolic blood pressure ≥130 mm Hg and a diastolic blood pressure ≥85 mm Hg,

(e) a fasting blood glucose level $\geq 110 \text{ mg/dL}$ or $\geq 100 \text{ mg/dL}$. It is assumed that patients diagnosed with impaired glucose tolerance (IGT), impaired fasting blood glucose (IFG), with insulin resistance and/or with metabolic syndrome suffer from an increased risk of developing a cardiovascular disease, such as for example myocardial infarction, coronary heart disease, heart insufficiency, thromboembolic events. A

glycemic control according to this invention may result in a reduction of the cardiovascular risks. Furthermore, the pharmaceutical composition and the

methods according to this invention are particularly suitable particular those patients who are diagnosed having one or

more of the following conditions (a) a higher age, in particular above 50 years,

- (b) male gender;
- (c) overweight, obesity (including class I, II and/or III 45 obesity), visceral obesity and/or abdominal obesity,
- (d) pre-transplant diabetes.
- (e) immunosuppression therapy.

A pharmaceutical composition or combination according to this invention, in particular due to the DPP-4 inhibitor 50 therein, exhibits a good safety profile. Therefore, a treatment or prophylaxis according to this invention is possible in those patients for which the mono-therapy with another antidiabetic drug, such as for example metformin, is contraindicated and/or who have an intolerance against such 55 drugs at therapeutic doses. In particular, a treatment or prophylaxis according to this invention may be advantageously possible in those patients showing or having an increased risk for one or more of the following disorders: renal insufficiency or diseases, cardiac diseases, cardiac 60 failure, hepatic diseases, pulmonal diseases, catabolytic states and/or danger of lactate acidosis, or female patients being pregnant or during lactation.

Furthermore, it can be found that the administration of a pharmaceutical composition or combination according to 65 this invention results in no risk or in a low risk of hypoglycemia. Therefore, a treatment or prophylaxis according to

42

this invention is also advantageously possible in those patients showing or having an increased risk for hypogly-

A pharmaceutical composition or combination according to this invention is particularly suitable in the long term treatment or prophylaxis of the diseases and/or conditions as described hereinbefore and hereinafter, in particular in the long term glycemic control in patients with type 2 diabetes mellitus.

The term "long term" as used hereinbefore and hereinafter indicates a treatment of or administration in a patient within a period of time longer than 12 weeks, preferably longer than 25 weeks, even more preferably longer than 1 year.

Therefore, a particular embodiment of the present invention provides a method for therapy, preferably oral therapy, for improvement, especially long term improvement, of glycemic control in patients with type 2 diabetes mellitus, especially in patients with late stage type 2 diabetes mellitus, 20 in particular in patients additionally diagnosed of overweight, obesity (including class I, class II and/or class III obesity), visceral obesity and/or abdominal obesity.

The effects mentioned above are observed both, when the DPP-4 inhibitor and the second and, optionally, third antidi-25 abetic agent are administered together, for example simultaneously in one single or two or three separate formulations, and/or when they are administered in alternation, for example successively in two or three separate formulations.

Within this invention it is to be understood that combinations or combined uses envisage the separate, sequential, simultaneous, concurrent, chronologically staggered or alternating administration of the components. It will be appreciated that the DPP-4 inhibitor and the other active substance(s) can be administered in a single dosage form or each in separate dosage forms.

In this context, "combination" or "combined" within the meaning of this invention also includes, without being limited, fixed and non-fixed forms and uses.

It will be appreciated that the amount of the pharmaceuin the treatment of patients after organ transplantation, in 40 tical composition according to this invention to be administered to the patient and required for use in treatment or prophylaxis according to the present invention will vary with the route of administration, the nature and severity of the condition for which treatment or prophylaxis is required, the age, weight and condition of the patient, concomitant medication and will be ultimately at the discretion of the attendant physician. In general, however, the DPP-4 inhibitor and, optionally, the second and/or third antidiabetic agent according to this invention are included in the pharmaceutical composition, combination or dosage form in an amount sufficient that by their administration the glycemic control in the patient to be treated is improved.

> In the following preferred ranges of the amount of the DPP-4 inhibitor, the second and/or third antidiabetic agent to be employed in the pharmaceutical composition and the methods and uses according to this invention are described. These ranges refer to the amounts to be administered per day with respect to an adult patient, in particular to a human being, for example of approximately 70 kg body weight, and can be adapted accordingly with regard to an administration 2, 3, 4 or more times daily and with regard to other routes of administration and with regard to the age of the patient. The ranges of the dosage and amounts are calculated for the individual active moiety. Advantageously, the combination therapy of the present invention utilizes lower dosages of the individual DPP-4 inhibitor and/or of the individual second and/or third antidiabetic agent used in monotherapy or used

in conventional therapeutics, thus avoiding possible toxicity and adverse side effects incurred when those agents are used as monotherapies

Within the scope of the present invention, the pharmaceutical composition or combination is preferably administered orally. Other forms of administration are possible and described hereinafter. Preferably the one or more dosage forms comprising the DPP-4 inhibitor and/or the second and/or the third antidiabetic agent is oral or usually well known.

In general, the amount of the DPP-4 inhibitor in the combinations, combination methods or combined uses of this invention is preferably in the range from 1/5 to 1/1 of the amount usually recommended for a monotherapy using said DPP-4 inhibitor.

A preferred dosage range of linagliptin when administered orally is 0.5 mg to 10 mg per day, preferably 2.5 mg to 10 mg, most preferably 1 mg to 5 mg per day. The preferred range of amounts in the pharmaceutical composition is 0.5 to 10 mg, in particular 1 to 5 mg. Examples of particular 20 dosage strengths are 1, 2.5, 5 or 10 mg. The application of the active ingredient may occur up to three times a day, preferably one or two times a day. Suitable formulations for linagliptin may be those formulations disclosed in the application WO 2007/128724, the disclosure of which is incorporated herein in its entirety.

Typical dosage strengths of the dual fixed dose combination (tablet) of linagliptin/metformin IR (immediate release) are 2.5/500 mg, 2.5/850 mg and 2.5/1000 mg, which may be administered 1-3 times a day, particularly twice a day.

Typical dosage strengths of the dual fixed dose combination (tablet) of linagliptin/metformin XR (extended release) are 5/500 mg, 5/1000 mg and 5/1500 mg, which may be administered 1-2 times a day, particularly once a day, preferably to be taken in the evening with meal, or 2.5/500 35 mg, 2.5/750 mg and 2.5/1000 mg, which may be administered 1-2 times a day, particularly once a day two tablets, preferably to be taken in the evening with meal.

A preferred dosage range of sitagliptin when administered orally is from 10 to 200 mg, in particular 25 to 150 mg per 40 day. A recommended dose of sitagliptin is 100 mg calculated for the active moiety (free base anhydrate) once daily or 50 mg twice daily. The preferred range of amounts in the pharmaceutical composition is 10 to 150 mg, in particular 25 to 100 mg. Examples are 25, 50, 75 or 100 mg. The 45 application of the active ingredient may occur up to three times a day, preferably one or two times a day. Equivalent amounts of salts of sitagliptin, in particular of the phosphate monohydrate can be calculated accordingly. Adjusted dosages of sitagliptin, for example 25 and 50 mg, are preferably 50 used for patients with renal failure.

A preferred dosage range of vildagliptin when administered orally is from 10 to 150 mg daily, in particular from 25 to 150 mg, 25 and 100 mg or 25 and 50 mg or 50 and 100 mg daily. For example the daily administration of vildagliptin is 50 or 100 mg. The preferred range of amounts in the pharmaceutical composition is 10 to 150 mg, in particular 25 to 100 mg. Examples are 25, 50, 75 or 100 mg. The application of the active ingredient may occur up to three times a day, preferably one or two times a day.

A preferred dosage range of alogliptin when administered orally is from 5 to 250 mg daily, in particular from 10 to 150 mg daily. The preferred range of amounts in the pharmaceutical composition is 5 to 150 mg, in particular 10 to 100 mg. Examples are 10, 12.5, 20, 25, 50, 75 and 100 mg. The 65 application of the active ingredient may occur up to three times a day, preferably one or two times a day.

44

A preferred dosage range of saxagliptin when administered orally is from 2.5 to 100 mg daily, in particular from 2.5 to 50 mg daily. The preferred range of amounts in the pharmaceutical composition is from 2.5 to 100 mg, in particular from 2.5 and 50 mg. Examples are 2.5, 5, 10, 15, 20, 30, 40, 50 and 100 mg. The application of the active ingredient may occur up to three times a day, preferably one or two times a day.

A preferred dosage range of dutogliptin when administered orally is from 50 to 400 mg daily, in particular from 100 to 400 mg daily. The preferred range of amounts in the pharmaceutical composition is from 50 to 400 mg. Examples are 50, 100, 200, 300 and 400 mg. The application of the active ingredient may occur up to three times a day, preferably one or two times a day.

A special embodiment of the DPP-4 inhibitors of this invention refers to those orally administered DPP-4 inhibitors which are therapeutically efficacious at low dose levels, e.g. at dose levels <100 mg or <70 mg per patient per day, preferably <50 mg, more preferably <30 mg or <20 mg, even more preferably from 1 mg to 10 mg (if required, divided into 1 to 4 single doses, particularly 1 or 2 single doses, which may be of the same size), particularly from 1 mg to 5 mg (more particularly 5 mg), per patient per day, preferentially, administered orally once-daily, more preferentially, at any time of day, administered with or without food. Thus, for example, the daily oral amount 5 mg BI 1356 can be given in a once daily dosing regimen (i.e. 5 mg BI 1356 once daily) or in a twice daily dosing regimen (i.e. 2.5 mg BI 1356 twice daily), at any time of day, with or without food.

In general, the amount of the second and/or third antidiabetic agent in the combinations, combination methods and/or combined uses of this invention is preferably in the range from 1/5 to 1/1 of the amount usually recommended for a monotherapy using said antidiabetic agent. Using lower dosages of the individual second and/or third antidiabetic agent compared with monotherapy could avoid or minimize possible toxicity and adverse side effects incurred when those agents are used as monotherapies.

A preferred dosage range of metformin when administered orally is 250 to 3000 mg, in particular 500 to 2000 mg per day. The preferred range of amounts in the pharmaceutical composition is 250 to 1000, in particular 500 to 1000 mg or 250 to 850 mg respectively. Examples are 500, 750, 850 or 1000 mg. Preferably the administration of said amounts is once, twice or three times daily. For example the amounts of 500, 750 and 850 mg preferably require oncedaily, twice-daily or three-times daily dosing and the amount of 1000 mg preferably requires once-daily or twice-daily dosing. Certain controlled or sustained release formulations allow a once-daily dosing. Metformin can be administered for example in the form as marketed under the trademarks GLUCOPHAGE-DTM or GLUCOPHAGE™, COPHAGE-XR™.

A preferred dosage range of pioglitazone when administered orally is 5 to 50 mg per day. The preferred range of amounts in the pharmaceutical composition is 5 to 50 mg, 10 to 45 mg and 15 to 45 mg respectively. Examples are 15, 30 or 45 mg. Preferably the administration of said amounts is once or twice daily, in particular once daily. Pioglitazone can be administered in the form as it is marketed for example under the trademark ACTOSTM.

A preferred dosage range of rosiglitazone when administered orally is 1 mg to 10 mg per day. The preferred range of amounts in the pharmaceutical composition is 1 to 10 mg, 2 to 8 mg, 4 to 8 mg and 1 to 4 mg. Examples are 1, 2, 4 or

8 mg. Preferably the administration of said amounts is once or twice daily. Preferably the dose should not exceed 8 mg daily. Rosiglitazone can be administered in the form as it is marketed for example under the trademark AVANDIATM.

A preferred dosage range of a thiazolidindione (other than 5 pioglitazone or rosiglitazone as described above) when administered orally is 2 to 100 mg per day. The preferred range of amounts in the pharmaceutical composition for an administration once, twice or three times daily is 2 to 100, 1 to 50 and 1 to 33 mg respectively.

A preferred dosage range of glibenclamide when administered orally is 0.5 to 15 mg, in particular 1 to 10 mg per day. The preferred range of amounts in the pharmaceutical composition is 0.5 to 5 mg, in particular 1 to 4 mg. Examples are 1.0, 1.75 and 3.5 mg. Preferably the administration of 15 said amounts is once, twice or three-times daily. Glibenclamide can be administered in the form as it is marketed for example under the trademark EUGLUCONTM.

A preferred dosage range of glimepiride when administered orally is 0.5 to 10 mg, in particular 1 to 6 mg per day. 20 The preferred range of amounts in the pharmaceutical composition is 0.5 to 10 mg, in particular 1 to 6 mg. Examples are 1, 2, 3, 4, and 6 mg. Preferably the administration of said amounts is once, twice or three-times daily, preferably once daily. Glimepiride can be administered in the form as it is 25 marketed for example under the trademark AMARYLTM.

A preferred dosage range of gliquidone when administered orally is 5 to 150 mg, in particular 15 to 120 mg per day. The preferred range of amounts in the pharmaceutical composition is 5 to 120 mg, in particular 5 to 30 mg. 30 Examples are 10, 20, 30 mg. Preferably the administration of said amounts is once, twice, three-times or four-times daily. Gliquidone can be administered in the form as it is marketed for example under the trademark GLURENORMTM.

A preferred dosage range of glibornuride when administered orally is 5 to 75 mg per day. The preferred range of amounts in the pharmaceutical composition is 5 to 75 mg, in particular 10 to 50 mg. Preferably the administration of said amounts is once, twice or three-times daily.

A preferred dosage range of gliclazide when administered orally is 20 to 300 mg, in particular 40 to 240 mg per day. The preferred range of amounts in the pharmaceutical composition is 20 to 240 mg, in particular 20 to 80 mg. Examples are 20, 30, 40 and 50 mg. Preferably the administration of 45 said amounts is once, twice or three-times daily.

A preferred dosage range of glisoxepide when administered orally is 1 to 20 mg, in particular 1 to 16 mg per day. The preferred range of amounts in the pharmaceutical composition is 1 to 8 mg, in particular 1 to 4 mg. Preferably the 50 administration of said amounts is once, twice, three-times or four-times daily.

A preferred dosage range of tolbutamide when administered orally is 100 to 3000 mg, preferably 500 to 2000 mg per day. The preferred range of amounts in the pharmaceutical composition is 100 to 1000 mg. Preferably the administration of said amounts is once or twice daily.

A preferred dosage range of glipizide when administered orally is 1 to 50 mg, in particular 2.5 to 40 mg per day. The preferred range of amounts in the pharmaceutical composition for an administration once, twice or three times daily is 1 to 50, 0.5 to 25 and 0.3 to 17 mg respectively.

A preferred dosage range of nateglinide when administered orally is 30 to 500 mg, in particular 60 to 360 mg per day. The preferred range of amounts in the pharmaceutical 65 composition is 30 to 120 mg. Examples are 30, 60 and 120 mg. Preferably the administration of said amounts is once,

46

twice or three-times daily. Nateglinide can be administered in the form as it is marketed for example under the trademark $STARLIX^{TM}$.

A preferred dosage range of repaglinide when administered orally is 0.1 to 16 mg, in particular 0.5 to 6 mg per day.

The preferred range of amounts in the pharmaceutical composition is 0.5 to 4 mg. Examples are 0.5, 1, 2 or 4 mg. Preferably the administration of said amounts is once, twice, three-times or four-times daily. Repaglinide can be administered in the form as it is marketed for example under the trademark NOVONORMTM.

A preferred dosage range of acarbose when administered orally is 50 to 1000 mg, in particular 50 to 600 mg per day. The preferred range of amounts in the pharmaceutical composition is 50 to 150 mg. Examples are 50 and 100 mg. Preferably the administration of said amounts is once, twice, three-times or four-times daily. Acarbose can be administered in the form as it is marketed for example under the trademark GlucobayTM.

A preferred dosage range of voglibose when administered orally is 100 to 1000 mg, in particular 200 to 600 mg per day. The preferred range of amounts in the pharmaceutical composition is 50 to 300 mg. Examples are 50, 100, 150, 200 and 300 mg. Preferably the administration of said amounts is once, twice, three-times or four-times daily. Voglibose can be administered in the form as it is marketed for example under the trademark BasenTM or VoglisanTM.

A preferred dosage range of miglitol when administered orally is 25 to 500 mg, in particular 25 to 300 mg per day.

The preferred range of amounts in the pharmaceutical composition is 25 to 100 mg. Examples are 25, 50 and 100 mg. Preferably the administration of said amounts is once, twice, three-times or four-times daily. Miglitol can be administered in the form as it is marketed for example under the trademark GlysetTM.

A preferred dosage range of GLP-1 analogues, in particular of exenatide is 5 to 30 μg, in particular 5 to 20 μg per day. The preferred range of amounts in the pharmaceutical composition is 5 to 10 μg. Examples are 5 and 10 μg. Preferably the administration of said amounts is once, twice, three-times or four-times daily by subcutaneous injection. Exenatide can be administered in the form as it is marketed for example under the trademark ByettaTM A long acting formulation, preferably for a once weekly subcutaneous injection, comprises an amount from 0.1 to 3.0 mg, preferably 0.5 to 2.0 mg exenatide. Examples are 0.8 mg and 2.0 mg. An example of a long acting formulation of exenatide is Byetta LARTM.

A preferred dosage range of liraglutide is 0.5 to 3 mg, in particular 0.5 to 2 mg per day. The preferred range of amounts in the pharmaceutical composition is 0.5 to 2 mg. Examples are 0.6, 1.2 and 1.8 mg. Preferably the administration of said amounts is once or twice daily by subcutaneous injection.

The amount of the DPP-4 inhibitor and the second and/or third therapeutic agent in the pharmaceutical composition and in the methods and uses of this invention correspond to the respective dosage ranges as provided hereinbefore. For example, preferred dosage ranges in a pharmaceutical composition, combination, method and use according to this invention are an amount of 0.5 to 10 mg (in particular 1 to 5 mg, especially 2.5 mg or 5 mg) of linagliptin and/or, optionally, an amount of 250 to 1000 mg (especially 500 mg, 850 mg or 1000 mg) of metformin. An oral administration once or twice daily is preferred.

In the combination methods and combined uses according to the present invention the DPP-4 inhibitor and the second

and/or third therapeutic agent are administered in combination including, without being limited, the active ingredients are administered at the same time, i.e. simultaneously, or essentially at the same time, or the active ingredients are administered in alternation, i.e. that at first one or two active ingredients are administered and after a period of time the other two or one active ingredients are administered, i.e. at least two of the three active ingredients are administered sequentially. The period of time may be in the range from 30 min to 12 hours. The administration which is in combination or in alternation may be once, twice, three times or four times daily, preferably once or twice daily.

With regard to combined administration of the DPP-4 inhibitor and the second and/or third antidiabetic agent, all three active ingredients may be present in one single dosage form, for example in one tablet or capsule, or one or two of the active ingredients may be present in a separate dosage form, for example in two different or identical dosage forms.

With regard to their administration in alternation, one or 20 two of the active ingredients are present in a separate dosage form, for example in two different or identical dosage forms. Therefore, a pharmaceutical combination of this invention may be present as single dosage forms which comprise the DPP-4 inhibitor and the second and, optionally, the third 25 antidiabetic agent. Alternatively a pharmaceutical combination of this invention may be present as two separate dosage forms wherein one dosage form comprises the DPP-4 inhibitor and the other dosage form comprises the second plus, optionally, the third antidiabetic agent, or, in case of a triple combination, one dosage form comprises the DPP-4 inhibitor plus either the second or the third antidiabetic agent and the other dosage form comprises the third or the second antidiabetic agent, respectively. Alternatively, in case of a triple combination, a pharmaceutical combination of this invention may be present as three separate dosage forms wherein one dosage form comprises the DPP-4 inhibitor and a second dosage form comprises the second antidiabetic agent and the third dosage form comprises the third antidiabetic agent. Alternatively, in case of a dual combination, a pharmaceutical combination of this invention may be present as two separate dosage forms wherein one dosage form comprises the DPP-4 inhibitor and the second dosage form comprises the second antidiabetic agent.

The case may arise in which an active ingredient has to be administered more often, for example twice per day, than the other active ingredient(s), which for example needs administration once daily. Therefore "administration in combination" also includes an administration scheme in which first 50 all active ingredients are administered in combination and after a period of time an active ingredient is administered again or vice versa.

Therefore, the present invention also includes pharmaceutical combinations which are present in separate dosage 55 forms wherein one dosage form comprises the DPP-4 inhibitor and the second and, optionally, the third, therapeutic agent and the other dosage form comprises the second and/or the third therapeutic agent only.

Thus, the present invention also includes pharmaceutical 60 compositions or combinations for separate, sequential, simultaneous, concurrent, alternate or chronologically staggered use of the active ingredients or components.

A pharmaceutical composition which is present as a separate or multiple dosage form, preferably as a kit of parts, 65 is useful in combination therapy to flexibly suit the individual therapeutic needs of the patient.

48

According to a first embodiment a kit of parts comprises (a) a first containment containing a dosage form comprising the DPP-4 inhibitor and at least one pharmaceutically acceptable carrier, and

- (b) a second containment containing a dosage form comprising the second antidiabetic agent and at least one pharmaceutically acceptable carrier, and, optionally,
- (c) a third containment containing a dosage form comprising the third antidiabetic agent and at least one pharmaceutically acceptable carrier.

According to a second embodiment a kit of parts comprises

- (a) a first containment containing a dosage form comprising the DPP-4 inhibitor and the second or third antidiabetic agent and at least one pharmaceutically acceptable carrier, and
- (b) a second containment containing a dosage form comprising the third or second antidiabetic agent, respectively, and at least one pharmaceutically acceptable carrier.

According to a third embodiment a kit of parts comprises (a) a first containment containing a dosage form comprising the DPP-4 inhibitor and at least one pharmaceutically acceptable carrier, and

(b) a second containment containing a dosage form comprising the second and third antidiabetic agent and at least one pharmaceutically acceptable carrier.

A further aspect of the present invention is a manufacture comprising the pharmaceutical combination being present as separate dosage forms according to the present invention and a label or package insert comprising instructions that the separate dosage forms are to be administered in combination.

According to a first embodiment a manufacture comprises (a) a pharmaceutical composition comprising a DPP-4 inhibitor according to the present invention and (b) a label or package insert which comprises instructions that the medicament may or is to be administered, for example in combination, with a medicament comprising a second antidiabetic agent according to the present invention or with a fixed or free combination (e.g. a medicament) comprising a second antidiabetic agent and a third antidiabetic agent according to the present invention.

According to a second embodiment a manufacture comprises (a) a second antidiabetic agent according to the present invention and (b) a label or package insert which comprises instructions that the medicament may or is to be administered, for example in combination, with a medicament comprising a DPP-4 inhibitor according to the present invention or with a a fixed or free-combination (e.g. a medicament) comprising a DPP-4 inhibitor and a third antidiabetic agent according to the present invention.

According to a third embodiment a manufacture comprises (a) a pharmaceutical composition comprising a DPP-4 inhibitor and a second antidiabetic agent according to the present invention and (b) a label or package insert which comprises instructions that the medicament may or is to be administered, for example in combination, with a medicament comprising a third antidiabetic agent according to the present invention.

The desired dose of the pharmaceutical composition according to this invention may conveniently be presented in a once daily or as divided dose administered at appropriate intervals, for example as two, three or more doses per day.

The pharmaceutical composition may be formulated for oral, rectal, nasal, topical (including buccal and sublingual), transdermal, vaginal or parenteral (including intramuscular, sub-cutaneous and intravenous) administration in liquid or

solid form or in a form suitable for administration by inhalation or insufflation. Oral administration is preferred. The formulations may, where appropriate, be conveniently presented in discrete dosage units and may be prepared by any of the methods well known in the art of pharmacy. All 5 methods include the step of bringing into association the active ingredient with one or more pharmaceutically acceptable carriers, like liquid carriers or finely divided solid carriers or both, and then, if necessary, shaping the product into the desired formulation.

The pharmaceutical composition may be formulated in the form of tablets, granules, fine granules, powders, capsules, caplets, soft capsules, pills, oral solutions, syrups, dry syrups, chewable tablets, troches, effervescent tablets, drops, suspension, fast dissolving tablets, oral fast-dispersing tablets, etc.

The pharmaceutical composition and the dosage forms preferably comprises one or more pharmaceutical acceptable carriers. Preferred carriers must be "acceptable" in the sense of being compatible with the other ingredients of the 20 formulation and not deleterious to the recipient thereof. Examples of pharmaceutically acceptable carriers are known to the one skilled in the art.

Pharmaceutical compositions suitable for oral administration may conveniently be presented as discrete units such 25 as capsules, including soft gelatin capsules, cachets or tablets each containing a predetermined amount of the active ingredient; as a powder or granules; as a solution, a suspension or as an emulsion, for example as syrups, elixirs or self-emulsifying delivery systems (SEDDS). The active 30 ingredients may also be presented as a bolus, electuary or paste. Tablets and capsules for oral administration may contain conventional excipients such as binding agents, fillers, lubricants, disintegrants, or wetting agents. The tablets may be coated according to methods well known in the 35 art. Oral liquid preparations may be in the form of, for example, aqueous or oily suspensions, solutions, emulsions, syrups or elixirs, or may be presented as a dry product for constitution with water or other suitable vehicle before use. Such liquid preparations may contain conventional additives 40 such as suspending agents, emulsifying agents, non-aqueous vehicles (which may include edible oils), or preservatives.

The pharmaceutical composition according to the invention may also be formulated for parenteral administration (e.g. by injection, for example bolus injection or continuous 45 infusion) and may be presented in unit dose form in ampoules, pre-filled syringes, small volume infusion or in multi-dose containers with an added preservative. The compositions may take such forms as suspensions, solutions, or emulsions in oily or aqueous vehicles, and may contain 50 formulatory agents such as suspending, stabilizing and/or dispersing agents. Alternatively, the active ingredients may be in powder form, obtained by aseptic isolation of sterile solid or by lyophilisation from solution, for constitution with a suitable vehicle, e.g. sterile, pyrogen-free water, before 55

Pharmaceutical compositions suitable for rectal administration wherein the carrier is a solid are most preferably presented as unit dose suppositories. Suitable carriers include cocoa butter and other materials commonly used in 60 the art, and the suppositories may be conveniently formed by admixture of the active compound(s) with the softened or melted carrier(s) followed by chilling and shaping in moulds.

For pharmaceutical application in warm-blooded verte- 65 brates, particularly humans, the compounds of this invention are usually used in dosages from 0.001 to 100 mg/kg body

50

weight, preferably at 0.1-15 mg/kg, in each case 1 to 4 times a day. For this purpose, the compounds, optionally combined with other active substances, may be incorporated together with one or more inert conventional carriers and/or diluents, e.g. with corn starch, lactose, glucose, microcrystalline cellulose, magnesium stearate, polyvinylpyrrolidone, citric acid, tartaric acid, water, water/ethanol, water/glycerol, water/sorbitol, water/polyethylene glycol, propylene glycol, cetylstearyl alcohol, carboxymethylcellulose or fatty substances such as hard fat or suitable mixtures thereof into conventional galenic preparations such as plain or coated tablets, capsules, powders, suspensions or suppositories.

The pharmaceutical compositions according to this invention comprising the DPP-4 inhibitors as defined herein are thus prepared by the skilled person using pharmaceutically acceptable formulation excipients as described in the art. Examples of such excipients include, without being restricted to diluents, binders, carriers, fillers, lubricants, flow promoters, crystallisation retardants, disintegrants, solubilizers, colorants, pH regulators, surfactants and emulsifiers.

Examples of suitable diluents for compounds according to embodiment A include cellulose powder, calcium hydrogen phosphate, erythritol, low substituted hydroxypropyl cellulose, mannitol, pregelatinized starch or xylitol. Among those diluents mannitol, low substituted hydroxypropyl cellulose and pregelatinized starch are to be emphasized.

Examples of suitable lubricants for compounds according to embodiment A include talc, polyethyleneglycol, calcium behenate, calcium stearate, hydrogenated castor oil or magnesium stearate. Among those lubricants magnesium stearate is to be emphasized.

Examples of suitable binders for compounds according to embodiment A include copovidone (copolymerisates of vinylpyrrolidon with other vinylderivates), hydroxypropyl methylcellulose (HPMC), hydroxypropylcellulose (HPC), polyvinylpyrrolidon (povidone), pregelatinized starch, or low-substituted hydroxypropylcellulose (L-HPC). Among those binders copovidone and pregelatinized starch are to be emphasized.

Examples of suitable disintegrants for compounds according to embodiment A include corn starch or crospovidone. Among those disintegrants corn starch is to be emphasized.

Suitable methods of preparing pharmaceutical formulations of the DPP-4 inhibitors according to embodiment A of the invention are

direct tabletting of the active substance in powder mixtures with suitable tabletting excipients;

granulation with suitable excipients and subsequent mixing with suitable excipients and subsequent tabletting as well as film coating; or

packing of powder mixtures or granules into capsules. Suitable granulation methods are

wet granulation in the intensive mixer followed by fluidised bed drying;

one-pot granulation;

fluidised bed granulation; or

dry granulation (e.g. by roller compaction) with suitable excipients and subsequent tabletting or packing into capsules.

An exemplary composition of a DPP-4 inhibitor according to embodiment A of the invention comprises the first diluent mannitol, pregelatinized starch as a second diluent with additional binder properties, the binder copovidone, the disintegrant corn starch, and magnesium stearate as lubricant; wherein copovidone and/or corn starch may be optional.

For details on dosage forms, formulations and administration of DPP-4 inhibitors of this invention, reference is made to scientific literature and/or published patent documents, particularly to those cited herein.

The pharmaceutical compositions (or formulations) may 5 be packaged in a variety of ways. Generally, an article for distribution includes a container that contains the pharmaceutical composition in an appropriate form. Tablets are typically packed in an appropriate primary package for easy handling, distribution and storage and for assurance of 10 proper stability of the composition at prolonged contact with the environment during storage. Primary containers for tablets may be bottles or blister packs.

A suitable bottle, e.g. for a pharmaceutical composition or combination comprising a DPP-4 inhibitor according to 15 embodiment A of the invention, may be made from glass or polymer (preferably polypropylene (PP) or high density polyethylene (HD-PE)) and sealed with a screw cap. The screw cap may be provided with a child resistant safety closure (e.g. press-and-twist closure) for preventing or ham- 20 pering access to the contents by children. If required (e.g. in regions with high humidity), by the additional use of a desiccant (such as e.g. bentonite clay, molecular sieves, or, preferably, silica gel) the shelf life of the packaged composition can be prolonged.

A suitable blister pack, e.g. for a pharmaceutical composition or combination comprising a DPP-4 inhibitor according to embodiment A of the invention, comprises or is formed of a top foil (which is breachable by the tablets) and a bottom part (which contains pockets for the tablets). The 30 top foil may contain a metallic foil, particularly an aluminium or aluminium alloy foil (e.g. having a thickness of 20 μm to 45 μm, preferably 20 μm to 25 μm) that is coated with a heat-sealing polymer layer on its inner side (sealing side). The bottom part may contain a multi-layer polymer 35 foil (such as e.g. poly(vinyl chloride) (PVC) coated with poly(vinylidene chloride) (PVDC); or a PVC foil laminated with poly(chlorotrifluoroethylene) (PCTFE)) or a multilayer polymer-metal-polymer foil (such as e.g. a coldtion).

The article may further comprise a label or package insert, which refer to instructions customarily included in commercial packages of therapeutic products, that may contain information about the indications, usage, dosage, adminis- 45 tration, contraindications and/or warnings concerning the use of such therapeutic products. In one embodiment, the label or package inserts indicates that the composition can be used for any of the purposes described herein.

The pharmaceutical compositions and methods according 50 to this invention show advantageous effects in the treatment and prevention of those diseases and conditions as described hereinbefore. The dual combinations show advantageous effects compared with monotherapy with an active ingredient. The triple combinations show advantageous effects 55 compared with dual therapy with one or two of the three active ingredients. Advantageous effects may be seen for example with respect to efficacy, dosage strength, dosage frequency, pharmacodynamic properties, pharmacokinetic properties, fewer adverse effects, convenience, compliance, 60

With respect to linagliptin, the methods of synthesis are known to the skilled person and as described in the literature, in particular as described in WO 2002/068420, WO 2004/018468, or WO 2006/048427, the disclosures of which 65 are incorporated herein. Polymorphous crystal modifications and formulations of particular DPP-4 inhibitors are dis**52**

closed in WO 2007/128721 and WO 2007/128724, respectively, the disclosures of which are incorporated herein in their entireties. Formulations of particular DPP-4 inhibitors with metformin or other combination partners are described in WO 2009/121945, the disclosure of which is incorporated herein in its entirety.

The methods of synthesis for the further DPP-4 inhibitors are described in the scientific literature and/or in published patent documents, particularly in those cited hereinbefore. The active ingredients, in particular the DPP-4 inhibitor and/or the second and/or the third antidiabetic agent, may be present in the form of a pharmaceutically acceptable salt. Pharmaceutically acceptable salts include, without being restricted thereto, such as salts of inorganic acid like hydrochloric acid, sulfuric acid and phosphoric acid; salts of organic carboxylic acid like oxalic acid, acetic acid, citric acid, malic acid, benzoic acid, maleic acid, fumaric acid, tartaric acid, succinic acid and glutamic acid and salts of organic sulfonic acid like methanesulfonic acid and p-toluenesulfonic acid. The salts can be formed by combining the compound and an acid in the appropriate amount and ratio in a solvent and decomposer. They can be also obtained by the cation or anion exchange from the form of other salts.

The active ingredients or a pharmaceutically acceptable salt thereof may be present in the form of a solvate such as a hydrate or alcohol adduct.

As different metabolic functional disorders often occur simultaneously, it is quite often indicated to combine a number of different active principles with one another. Thus, depending on the functional disorders diagnosed, improved treatment outcomes may be obtained if a DPP-4 inhibitor is combined with active substances customary for the respective disorders, such as e.g. one or more active substances selected from among the other antidiabetic substances, especially active substances that lower the blood sugar level or the lipid level in the blood, raise the HDL level in the blood, lower blood pressure or are indicated in the treatment of atherosclerosis or obesity.

The DPP-4 inhibitors mentioned above—besides their use formable laminated PVC/aluminium/polyamide composi- 40 in mono-therapy—may also be used in conjunction with other active substances, by means of which improved treatment results can be obtained. Such a combined treatment may be given as a free combination of the substances or in the form of a fixed combination, for example in a tablet or capsule. Pharmaceutical formulations of the combination partner needed for this may either be obtained commercially as pharmaceutical compositions or may be formulated by the skilled man using conventional methods. The active substances which may be obtained commercially as pharmaceutical compositions are described in numerous places in the prior art, for example in the list of drugs that appears annually, the "Rote Liste®" of the federal association of the pharmaceutical industry, or in the annually updated compilation of manufacturers' information on prescription drugs known as the "Physicians' Desk Reference".

Examples of antidiabetic combination partners are metformin; sulphonylureas such as glibenclamide, tolbutamide, glimepiride, glipizide, gliquidon, glibornuride and gliclazide; nateglinide; repaglinide; thiazolidinediones such as rosiglitazone and pioglitazone; PPAR gamma modulators such as metaglidases; PPAR-gamma agonists such as rivoglitazone, mitoglitazone, INT-131 or balaglitazone; PPARgamma antagonists; PPAR-gamma/alpha modulators such as tesaglitazar, muraglitazar, aleglitazar, indeglitazar and KRP297; PPAR-gamma/alpha/delta modulators such as e.g. lobeglitazone; AMPK-activators such as AICAR; acetyl-CoA carboxylase (ACC1 and ACC2) inhibitors; diacylglyccell GCRP agonists such as SMT3-receptor-agonists and

GPR119, such as the GPR119 agonists 5-ethyl-2-{4-[4-(4tetrazol-1-yl-phenoxymethyl)-thiazol-2-yl]-piperidin-1-yl}pyrimidine or 5-[1-(3-isopropyl-[1,2,4]oxadiazol-5-yl)-pip-5 eridin-4-ylmethoxy]-2-(4-methanesulfonyl-phenyl)pyridine; 11β-HSD-inhibitors; FGF19 agonists analogues; alpha-glucosidase blockers such as acarbose, voglibose and miglitol; alpha2-antagonists; insulin and insulin analogues such as human insulin, insulin lispro, insulin glusilin, r-DNA-insulinaspart, NPH insulin, insulin detemir, insulin degludec, insulin tregopil, insulin zinc suspension and insulin glargin; Gastric inhibitory Peptide (GIP); amylin and amylin analogues (e.g. pramlintide or davalintide); 15 GLP-1 and GLP-1 analogues such as Exendin-4, e.g. exenatide, exenatide LAR, liraglutide, taspoglutide, lixisenatide (AVE-0010), LY-2428757, dulaglutide (LY-2189265), semaglutide or albiglutide; SGLT2-inhibitors such as e.g. flozin, ipragliflozin or tofogliflozin; inhibitors of protein tyrosine-phosphatase (e.g. trodusquemine); inhibitors of glucose-6-phosphatase; fructose-1,6-bisphosphatase modulators; glycogen phosphorylase modulators; glucagon recepphosphoenolpyruvatecarboxykinase 25 antagonists: (PEPCK) inhibitors; pyruvate dehydrogenasekinase (PDK) inhibitors; inhibitors of tyrosine-kinases (50 mg to 600 mg) such as PDGF-receptor-kinase (cf. EP-A-564409, WO 98/35958, U.S. Pat. No. 5,093,330, WO 2004/005281, and WO 2006/041976) or of serine/threonine kinases; glucokinase/regulatory protein modulators incl. glucokinase activators; glycogen synthase kinase inhibitors; inhibitors of the SH2-domain-containing inositol 5-phosphatase type 2 (SHIP2); IKK inhibitors such as high-dose salicylate; JNK1 inhibitors; protein kinase C-theta inhibitors; beta 3 agonists such as ritobegron, YM 178, solabegron, talibegron, N-5984, GRC-1087, rafabegron, FMP825; aldosereductase inhibitors such as AS 3201, zenarestat, fidarestat, epalrestat, ranirestat, NZ-314, CP-744809, and CT-112; SGLT-1 or 40 SGLT-2 inhibitors, such as e.g. dapagliflozin, sergliflozin, atigliflozin, canagliflozin or (1S)-1,5-anhydro-1-[3-(1-benzothiophen-2-ylmethyl)-4-fluorophenyl]-D-glucitol; KV 1.3 channel inhibitors; GPR40 modulators such as e.g. [(3S)-6-({2',6'-dimethyl-4'-[3-(methylsulfonyl)propoxy]biphenyl-3- 45 yl}methoxy)-2,3-dihydro-1-benzofuran-3-yl]acetic SCD-1 inhibitors; CCR-2 antagonists; dopamine receptor agonists (bromocriptine mesylate [Cycloset]); 4-(3-(2,6-di-

stimulants; and other DPP IV inhibitors. Metformin is usually given in doses varying from about 500 mg to 2000 mg up to 2500 mg per day using various dosing regimens from about 100 mg to 500 mg or 200 mg to 850 mg (1-3 times a day), or about 300 mg to 1000 mg once or twice a day, or delayed-release metformin in doses 55 of about 100 mg to 1000 mg or preferably 500 mg to 1000 mg once or twice a day or about 500 mg to 2000 mg once a day. Particular dosage strengths may be 250, 500, 625, 750, 850 and 1000 mg of metformin hydrochloride.

acid;

sirtuin

methylbenzyloxy)phenyl)-4-oxobutanoic

For children 10 to 16 years of age, the recommended 60 starting dose of metformin is 500 mg given once daily. If this dose fails to produce adequate results, the dose may be increased to 500 mg twice daily. Further increases may be made in increments of 500 mg weekly to a maximum daily dose of 2000 mg, given in divided doses (e.g. 2 or 3 divided 65 doses). Metformin may be administered with food to decrease nausea.

A dosage of pioglitazone is usually of about 1-10 mg, 15 mg, 30 mg, or 45 mg once a day.

Rosiglitazone is usually given in doses from 4 to 8 mg once (or divided twice) a day (typical dosage strengths are 2, 4 and 8 mg).

Glibenclamide (glyburide) is usually given in doses from 2.5-5 to 20 mg once (or divided twice) a day (typical dosage strengths are 1.25, 2.5 and 5 mg), or micronized glibenclamide in doses from 0.75-3 to 12 mg once (or divided twice) a day (typical dosage strengths are 1.5, 3, 4.5 and 6 mg).

Glipizide is usually given in doses from 2.5 to 10-20 mg once (or up to 40 mg divided twice) a day (typical dosage strengths are 5 and 10 mg), or extended-release glibenclamide in doses from 5 to 10 mg (up to 20 mg) once a day (typical dosage strengths are 2.5, 5 and 10 mg).

Glimepiride is usually given in doses from 1-2 to 4 mg (up to 8 mg) once a day (typical dosage strengths are 1, 2 and 4 mg).

A dual combination of glibenclamide/metformin is usudapagliflozin, sergliflozin (KGT-1251), atigliflozin, canagli- 20 ally given in doses from 1.25/250 once daily to 10/1000 mg twice daily. (typical dosage strengths are 1.25/250, 2.5/500 and 5/500 mg).

> A dual combination of glipizide/metformin is usually given in doses from 2.5/250 to 10/1000 mg twice daily (typical dosage strengths are 2.5/250, 2.5/500 and 5/500 mg).

A dual combination of glimepiride/metformin is usually given in doses from 1/250 to 4/1000 mg twice daily.

A dual combination of rosiglitazone/glimepiride is usually given in doses from 4/1 once or twice daily to 4/2 mg twice daily (typical dosage strengths are 4/1, 4/2, 4/4, 8/2 and 8/4 mg).

A dual combination of pioglitazone/glimepiride is usually given in doses from 30/2 to 30/4 mg once daily (typical 35 dosage strengths are 30/4 and 45/4 mg).

A dual combination of rosiglitazone/metformin is usually given in doses from 1/500 to 4/1000 mg twice daily (typical dosage strengths are 1/500, 2/500, 4/500, 2/1000 and 4/1000 mg).

A dual combination of pioglitazone/metformin is usually given in doses from 15/500 once or twice daily to 15/850 mg thrice daily (typical dosage strengths are 15/500 and 15/850 mg).

The non-sulphonylurea insulin secretagogue nateglinide is usually given in doses from 60 to 120 mg with meals (up to 360 mg/day, typical dosage strengths are 60 and 120 mg); repaglinide is usually given in doses from 0.5 to 4 mg with meals (up to 16 mg/day, typical dosage strengths are 0.5, 1 and 2 mg). A dual combination of repaglinide/metformin is 50 available in dosage strengths of 1/500 and 2/850 mg.

Acarbose is usually given in doses from 25 to 100 mg with meals. Miglitol is usually given in doses from 25 to 100 mg with meals.

Examples of combination partners that lower the lipid level in the blood are HMG-CoA-reductase inhibitors such as simvastatin, atorvastatin, lovastatin, fluvastatin, pravastatin, pitavastatin and rosuvastatin; fibrates such as bezafibrate, fenofibrate, clofibrate, gemfibrozil, etofibrate and etofyllinclofibrate; nicotinic acid and the derivatives thereof such as acipimox; PPAR-alpha agonists; PPAR-delta agonists; inhibitors of acyl-coenzyme A:cholesterolacyltransferase (ACAT; EC 2.3.1.26) such as avasimibe; cholesterol resorption inhibitors such as ezetimib; substances that bind to bile acid, such as cholestyramine, colestipol and colesevelam; inhibitors of bile acid transport; HDL modulating active substances such as D4F, reverse D4F, LXR modulating active substances and FXR modulating active sub-

stances; CETP inhibitors such as torcetrapib, JTT-705 (dalcetrapib) or compound 12 from WO 2007/005572 (anacetrapib); LDL receptor modulators; MTP inhibitors (e.g. lomitapide); and ApoB100 antisense RNA.

A dosage of atorvastatin is usually from 1 mg to 40 mg or 5 10 mg to 80 mg once a day.

Examples of combination partners that lower blood pressure are beta-blockers such as atenolol, bisoprolol, celiprolol, metoprolol and carvedilol; diuretics such as hydrochlortalidon, furosemide, chlorothiazide, xipamide, piretanide, torasemide, spironolactone, eplerenone, amiloride and triamterene; calcium channel blockers such as amlodipine, nifedipine, nitrendipine, nisoldipine, nicardipine, felodipine, lacidipine, lercanipidine, manidipine, isradipine, nilvadipine, verapamil, gallopamil and diltiazem; ACE inhibitors such as ramipril, lisinopril, cilazapril, quinapril, captopril, enalapril, benazepril, perindopril, fosinopril and trandolapril; as well as angiotensin II receptor losartan, irbesartan, olmesartan, azilsartan and eprosartan.

A dosage of telmisartan is usually from 20 mg to 320 mg or 40 mg to 160 mg per day.

Examples of combination partners which increase the HDL level in the blood are Cholesteryl Ester Transfer 25 Protein (CETP) inhibitors; inhibitors of endothelial lipase; regulators of ABC1; LXRalpha antagonists; LXRbeta agonists; PPAR-delta agonists; LXRalpha/beta regulators, and substances that increase the expression and/or plasma concentration of apolipoprotein A-I.

Examples of combination partners for the treatment of obesity are sibutramine; tetrahydrolipstatin (orlistat); alizyme (cetilistat); dexfenfluramine; axokine; cannabinoid receptor 1 antagonists such as the CB1 antagonist 35 rimonobant; MCH-1 receptor antagonists; MC4 receptor agonists; NPY5 as well as NPY2 antagonists (e.g. velneperit); beta3-AR agonists such as SB-418790 and AD-9677; 5HT2c receptor agonists such as APD 356 (lorcaserin); myostatin inhibitors; Acrp30 and adiponectin; steroyl CoA 40 desaturase (SCD1) inhibitors; fatty acid synthase (FAS) inhibitors; CCK receptor agonists; Ghrelin receptor modulators; Pyy 3-36; orexin receptor antagonists; and tesofensine; as well as the dual combinations bupropion/naltrexone, bupropion/zonisamide, topiramate/phentermine 45 and pramlintide/metreleptin.

Examples of combination partners for the treatment of atherosclerosis are phospholipase A2 inhibitors; inhibitors of tyrosine-kinases (50 mg to 600 mg) such as PDGFreceptor-kinase (cf. EP-A-564409, WO 98/35958, U.S. Pat. 50 No. 5,093,330, WO 2004/005281, and WO 2006/041976); oxLDL antibodies and oxLDL vaccines; apoA-1 Milano; ASA; and VCAM-1 inhibitors.

The present invention is not to be limited in scope by the specific embodiments described herein. Various modifications of the invention in addition to those described herein may become apparent to those skilled in the art from the present disclosure. Such modifications are intended to fall within the scope of the appended claims.

All patent applications cited herein are hereby incorporated by reference in their entireties.

Further embodiments, features and advantages of the present invention may become apparent from the following examples. The following examples serve to illustrate, by 65 way of example, the principles of the invention without restricting it.

56

EXAMPLES

Example 1

BI 1356, a Potent and Selective DPP-4 Inhibitor, is Safe and Efficacious in Patients with Inadequately Controlled Type 2 Diabetes Despite Metformin Therapy

Efficacy and safety of BI 1356 (1, 5, or 10 mg qd), a potent and selective dipeptidyl peptidase-4 (DPP-4) inhibitor, was examined in inadequately controlled, metformin-treated (MET, ≥1 g daily) type 2 diabetic patients (T2DM; HbA1c at baseline 7.5-10.0%). Effects were compared to add-on of placebo (PBO) or of open label glimepiride (GLIM; 1 to 3 mg qd) in a 12-week randomized, double-blind study. Antidiabetic medication other than metformin was washed out for 6 weeks (34.7% of the patients).

The primary endpoint was change from baseline in blockers (ARBs) such as telmisartan, candesartan, valsartan, 20 HbA1c, adjusted for prior antidiabetic medication. 333 patients (mean baseline HbA1c 8.3%; fasting plasma glucose [FPG] 185 mg/dL) were randomized to BI 1356, PBO or open-label GLIM. After 12 weeks, BI 1356 treatment resulted in significant placebo corrected mean reductions in HbA1c (BI 1356 1 mg, n=65, -0.39%; 5 mg, n=66, -0.75%; 10 mg, n=66, -0.73%). Patients receiving GLIM demonstrated a slightly greater mean PBO corrected reduction in HbA1c at Week 12 (n=64, -0.90%). Reductions in FPG from baseline to Week 12 with BI 1356 were statistically significant (1 mg, -19 mg/dL; 5 mg, -35 mg/dL; 10 mg, -30 mg/dL). Hence, a dose-response relationship was demonstrated for HbA1c and FPG, reaching an effect plateau at 5 mg of BI 1356. For this dose, >80% DPP-4 inhibition at trough in >80% of the patients at week 12 was achieved.

> In total, 106 patients (43.1%) experienced adverse events (AEs) with similar incidences across all treatments. Most frequently reported episodes were nasopharyngitis (7.5%), diarrhoea (3.3%), and nausea (3.0%). Drug-related hypoglycaemia did not occur with BI 1356 or PBO but in 3 patients receiving GLIM. Ten patients (3.7%) experienced serious AEs but none of these events were considered drug-related.

> The addition of BI 1356 to MET in patients with T2DM inadequately controlled on MET alone achieved clinically relevant and statistically significant reductions in HbA1c. Combination treatment with BI 1356 1, 5, and 10 mg and MET was well tolerated and no case of hypoglycaemia was reported. The incidence of AEs was comparable with BI 1356 and PBO.

Example 2

The usability of a DPP-4 inhibitor or combination according to this invention for the purpose of the present invention (e.g. the beneficial effect on glycemic control) can be tested using clinical trials.

For example, in a randomised, double-blind, placebocontrolled, parallel group trial, the safety and efficacy of a DPP-4 inhibitor according to the invention (e.g. 5 mg of 60 linagliptin administered orally once daily) is tested in patients with type 2 diabetes with insufficient glycemic control (HbA1c from 7.0% to 10% or from 7.5% to 10% or from 7.5% to 11%) despite a therapy with one or two conventional antihyperglycemic agents, e.g. selected from metformin, thiazolidindiones (e.g. pioglitazone), sulfonylureas, glinides, inhibitors of alpha-glucosidase, GLP-1 or GLP-1 analogues, and insulin or insulin analogues.

57

In the study with the sulphonylurea drug the efficacy and safety of a DPP-4 inhibitor according to this invention versus placebo added to a background therapy of a sulphonylurea is investigated (2 week placebo run-in phase; 18 weeks double-blind treatment followed by 1 week follow up after study medication termination; background therapy with a sulphonylurea drug is administered throughout the entire trial duration, including placebo run-in phase, in an unchanged dosage).

The success of the treatment is tested by determining the HbA1c value, by comparison with the initial value and/or with the value of the placebo group. A significant change in the HbA1c value compared with the initial value and/or the placebo value demonstrates the efficacy of the DPP-4 inhibitor for the treatment. The success of the treatment can be also tested by determining the fasting plasma glucose values, by comparison with the initial values and/or with the values of the placebo group. A significant drop in the fasting glucose levels demonstrates the efficacy of the treatment. Also, the occurrence of a treat to target response (i.e. an HbA1c under treatment <7%) demonstrates the efficacy of the treatment.

The safety and tolerability of the treatment is investigated by assessing patient's condition and relevant changes from baseline, e.g. incidence and intensity of adverse events (such ²⁵ as e.g. hypoglycaemic episodes or the like) or weight gain.

Example 3

Treatment of Pre-Diabetes

The efficacy of a pharmaceutical composition or combination according to the invention in the treatment of prediabetes characterised by pathological fasting glucose and/ or impaired glucose tolerance can be tested using clinical 35 studies. In studies over a shorter period (e.g. 2-4 weeks) the success of the treatment is examined by determining the fasting glucose values and/or the glucose values after a meal or after a loading test (oral glucose tolerance test or food tolerance test after a defined meal) after the end of the period 40 of therapy for the study and comparing them with the values before the start of the study and/or with those of a placebo group. In addition, the fructosamine value can be determined before and after therapy and compared with the initial value and/or the placebo value. A significant drop in the fasting or 45 non-fasting glucose levels demonstrates the efficacy of the treatment. In studies over a longer period (12 weeks or more) the success of the treatment is tested by determining the HbA1c value, by comparison with the initial value and/or with the value of the placebo group. A significant 50 change in the HbA1c value compared with the initial value and/or the placebo value demonstrates the efficacy of the DPP-4 inhibitors or combinations according to the present invention for treating pre-diabetes.

Example 4

Preventing Manifest Type 2 Diabetes

Treating patients with pathological fasting glucose and/or 60 impaired glucose tolerance (pre-diabetes) is also in pursuit of the goal of preventing the transition to manifest type 2 diabetes. The efficacy of a treatment can be investigated in a comparative clinical study in which pre-diabetes patients are treated over a lengthy period (e.g. 1-5 years) with either 65 a pharmaceutical composition or combination according to this invention or with placebo or with a non-drug therapy or

58

other medicaments. During and at the end of the therapy, by determining the fasting glucose and/or a loading test (e.g. oGTT), a check is made to determine how many patients exhibit manifest type 2 diabetes, i.e. a fasting glucose level of >125 mg/dl and/or a 2 h value according to oGTT of >199 mg/dl. A significant reduction in the number of patients who exhibit manifest type 2 diabetes when treated with a DPP-4 inhibitor or combination according to the present invention as compared to one of the other forms of treatment, demonstrates the efficacy in preventing a transition from prediabetes to manifest diabetes.

Example 5

Treatment of Type 2 Diabetes

Treating patients with type 2 diabetes with the pharmaceutical composition or combination according to the invention, in addition to producing an acute improvement in the glucose metabolic situation, prevents a deterioration in the metabolic situation in the long term. This can be observed is patients are treated for a longer period, e.g. 3 months to 1 year or even 1 to 6 years, with the pharmaceutical composition or combination according to the invention and are compared with patients who have been treated with other antidiabetic medicaments. There is evidence of therapeutic success compared with patients treated with other antidiabetic medicaments if no or only a slight increase in the fasting glucose and/or HbA1c value is observed. Further evidence of therapeutic success is obtained if a significantly smaller percentage of the patients treated with a pharmaceutical composition or combination according to the invention, compared with patients who have been treated with other medicaments, undergo a deterioration in the glucose metabolic position (e.g. an increase in the HbA1c value to >6.5% or >7%) to the point where treatment with an additional oral antidiabetic medicament or with insulin or with an insulin analogue is indicated.

Example 6

Treatment of Insulin Resistance

In clinical studies running for different lengths of time (e.g. 2 weeks to 12 months) the success of the treatment is checked using a hyperinsulinaemic euglycaemic glucose clamp study. A significant rise in the glucose infusion rate at the end of the study, compared with the initial value or compared with a placebo group, or a group given a different therapy, proves the efficacy of a DPP-4 inhibitor, pharmaceutical composition or combination according to the present invention according to the invention in the treatment of insulin resistance.

Example 7

Treatment of Hyperglycaemia

In clinical studies running for different lengths of time (e.g. 1 day to 24 months) the success of the treatment in patients with hyperglycaemia is checked by determining the fasting glucose or non-fasting glucose (e.g. after a meal or a loading test with oGTT or a defined meal). A significant fall in these glucose values during or at the end of the study, compared with the initial value or compared with a placebo group, or a group given a different therapy, proves the efficacy of a DPP-4 inhibitor, pharmaceutical composition or

combination according to the present invention according to the invention in the treatment of hyperglycaemia.

Example 8

Prevention of Micro- or Macrovascular Complications

The treatment of type 2 diabetes or pre-diabetes patients with a DPP-4 inhibitor, pharmaceutical composition or combination according to the invention prevents or reduces or reduces the risk of developing microvascular complications (e.g. diabetic neuropathy, diabetic retinopathy, diabetic 15 nephropathy, diabetic foot, diabetic ulcer) or macrovascular complications (e.g. myocardial infarct, acute coronary syndrome, unstable angina pectoris, stable angina pectoris, stroke, peripheral arterial occlusive disease, cardiomyopathy, heart failure, heart rhythm disorders, vascular resteno- 20 sis). Type 2 diabetes or patients with pre-diabetes are treated long-term, e.g. for 1-6 years, with a pharmaceutical composition or combination according to the invention and compared with patients who have been treated with other antidiabetic medicaments or with placebo. Evidence of the therapeutic success compared with patients who have been treated with other antidiabetic medicaments or with placebo can be found in the smaller number of single or multiple complications. In the case of macrovascular events, diabetic foot and/or diabetic ulcer, the numbers are counted by anamnesis and various test methods. In the case of diabetic retinopathy the success of the treatment is determined by computer-controlled illumination and evaluation of the background to the eye or other ophthalmic methods. In the 35 case of diabetic neuropathy, in addition to anamnesis and clinical examination, the nerve conduction rate can be measured using a calibrated tuning fork, for example. With regard to diabetic nephropathy the following parameters may be investigated before the start, during and at the end 40 of the study: secretion of albumin, creatinine clearance, serum creatinin values, time taken for the serum creatinine values to double, time taken until dialysis becomes necessary.

Example 9

Treatment of Metabolic Syndrome

The efficacy of a DPP-4 inhibitor, pharmaceutical composition or combination according to the present invention according to the invention can be tested in clinical studies with varying run times (e.g. 12 weeks to 6 years) by 55 determining the fasting glucose or non-fasting glucose (e.g. after a meal or a loading test with oGTT or a defined meal) or the HbA1c value. A significant fall in these glucose values or HbA1c values during or at the end of the study, compared with the initial value or compared with a placebo group, or 60 a group given a different therapy, proves the efficacy of an active substance or combination of active substances in the treatment of Metabolic Syndrome. Examples of this are a reduction in systolic and/or diastolic blood pressure, a lowering of the plasma triglycerides, a reduction in total or 65 LDL cholesterol, an increase in HDL cholesterol or a reduction in weight, either compared with the starting value

60

at the beginning of the study or in comparison with a group of patients treated with placebo or a different therapy.

Example 10

Therapeutic Response to DPP-4 Inhibitor Treatment

Genomic DNA samples from individual patients enrolled in a clinical trial (e.g. a clinical study as described herein) for a DPP-4 inhibitor (e.g. linagliptin, e.g. in a daily oral amount of 5 mg, optionally in combination with one or more other antidiabetic agents) are obtained and genotyped for variation(s) (e.g. polymorphisms) in one or more candidate genes selected from TCF7L2, KCNJ11, PPARG and GLP1R, particularly for a TCF7L2 risk genotype as described herein, and evaluated relative to each patients response in the clinical trial (cf., e.g., Example 21). The association between the likelihood (e.g., increased, decreased, or no likelihood) of a favorable DPP-4 inhibitor therapy response (e.g. favorable change in HbA1c value) and genetic variations (e.g. TCF7L2 risk genotypes) or references can be investigated by applying statistical analysis to the results of genotyping.

The probability of the likelihood of a favorable response of an individual resulting from treating said individual with the DPP-4 inhibitor may be thus determined by such genotyping a nucleic acid sample of the individual, for example by detecting one or more single nucleotide polymorphisms within the TCF7L2 gene, for example one SNP selected from rs7903146, rs12255372 and rs10885406, or by detecting the respective wild-type genotype (cf., e.g., Example 21).

Methods for genotyping, i.e. determining genetic variations (e.g. polymorphisms, particularly those described herein) from patients' nucleic acid samples are known in the art. For example, molecular genetic methods to detect single nucleotide polymorphisms, e.g. within the TCF7L2 gene, may be based on genetic sequencing, microarray or PCR analysis.

Example 11

Linagliptin Monotherapy Improves Glycemic Control and Measures of β -Cell Function in Type 2 Diabetes

In a multi-center, 24 week, randomized, double-blind, placebo-controlled, parallel group study, the effects of linagliptin (LI) monotherapy (5 mg qd) are compared with placebo (PBO) in drug naïve or previously treated patients (pts) with type 2 diabetes mellitus (T2DM) (baseline HbA1c 4.9-10.6%). Randomization to LI (n=336) or PBO (n=167) follows a 2-week PBO run-in (previously treated pts go without medication for 4 wks prior to this). Mean baseline demographics (HbA1c, 8.0% [SD 0.87]; fasting plasma glucose (FPG), 166.0 mg/dL [41.1]; body mass index (BMI), 29.05 kg/m2 [4.81]; age, 55.7 yrs [10.2]) are similar in both groups. The primary endpoint is the change from baseline in HbA1c after 24 wks of treatment. LI shows a PBO-adjusted change in HbA1c from baseline of -0.69% (p<0.0001) with a continuous HbA1c reduction over time of -0.46% at 6 weeks to -0.69% at 24 weeks (both p<0.0001). LI patients are >4-fold more likely to achieve a reduction in HbA1c of $\geq 0.5\%$ at 24 weeks than PBO (47.1% vs 19.0%; p<0.0001). For patients with baseline HbA1c ≥7.0% a significant greater number of LI-treated compared to PBO-treated patients achieve a target reduction of HbA1c to <7.0% at 24 weeks (25.2% vs. 11.6%; odds ratio of 2.9, p=0.0006).

Patients with baseline HbA1c levels of ≥9.0% show the greatest reduction in HbA1c (-0.86%) from baseline. FPG improves by -23.3 mg/dL (p<0.0001) vs. PBO. In a meal tolerance test, the LI patients show a greater reduction in the adjusted mean change from baseline at week 24 for 2-hr 5 postprandial glucose (PPG) (-58.4 mg/dL; p<0.0001) vs. PBO. LI improves insulin secretion (p<0.05), as shown by changes in HOMA-% B index (LI, 5.02 vs PBO, -17.2 [(mU/L)/(mmol/L)]), proinsulin/insulin ratio (LI, -0.015 vs PBO, 0.024) and the disposition index (LI, 3.05 vs PBO, 10 -0.68). The proportion of patients reporting at least one adverse event (AE) is similar for both groups (52.4% LI; 58.7% PBO). Hypoglycemia is rare, occurring in 1 patients in each of the groups. Serious AEs are reported in both groups (LI, 3.0%; PBO, 4.2%) but are not considered 15 comparable to placebo. drug-related. Linagliptin trough levels in patients with mild and moderate renal impairment are comparable to patients with normal renal function.

Conclusion: Linagliptin monotherapy shows a significant, clinically meaningful and sustained improvement in glyce- 20 mic control reflected in changes in FPG and HbA1c, and accompanied by β-cell function improvements. Linagliptin is safe and well tolerated with no clinically significant changes in body weight or waist circumference. Linagliptin impairment are comparable to patients with normal renal function, supporting that no dose adjustment is required in renally impaired patients.

Example 12

Efficacy and Safety of Linagliptin in Type 2 Diabetes Inadequately Controlled on Metformin Monotherapy

A multi-center, 24-week, randomized, placebo-controlled, double-blind, parallel group study examines the efficacy and safety of linagliptin (LI) administered as add-on therapy to metformin (MET) in type 2 diabetes mellitus (T2DM) hyperglycemic patients with insufficient glycemic control 40 (HbA1c≥7 to 510.0% for patients previously treated only with metformin, or ≥6.5 to ≤9.0% for patients previously treated with additional oral antihyperglycemic drugs). Subjects who enter the screening period discontinue previous antidiabetic medication other than MET (≥1500 mg/day) for 45 6 weeks (including a placebo (PBO) run-in period during the last 2 weeks) prior to randomization to LI (n=524) or PBO (n=177). Mean baseline characteristics and demographics (HbA1c, 8.1%; fasting plasma glucose [FPG], 168.8 mg/dL; age, 56.5 yrs; BMI, 29.9 kg/m2) are similar between groups. 50 The primary endpoint is the change from baseline HbA1c after 24 weeks of treatment, evaluated with an analysis of covariance (ANCOVA) adjusted for baseline HbA1c and prior antidiabetic medication. After 24 weeks of treatment, and PBO+MET is -0.64% (p<0.0001) in favor of LI+MET for change in HbA1c (%). Patients with a baseline HbA1c of ≥7.0% who receive LI+MET are more likely to achieve an HbA1c 57.0% relative to those receiving placebo+MET (26.2% vs. 9.2%, respectively; odds ratio, 4.4; p=0.0001). At 60 week 24 LI+MET is superior to PBO+MET in reducing the mean fasting plasma glucose (FPG) from baseline (-21.1 mg/dL; p<0.0001). At study-end, 2 hr post-prandial glucose (PPG) analyzed in meal tolerance tests shows a significantly greater (p<0.0001) mean reduction from baseline for the 65 LI+MET treated (-67.1 mg/dL) versus the PBO+MET group. The proportion of patients reporting at least one

62

adverse event (AE) is comparable within the LI+MET and PBO+MET groups (52.8% and 55.4%, respectively). Hypoglycemia is rare, occurring in 5 PBO+MET patients (2.8%) and 3 LI+MET patients (0.6%), all episodes being of mild intensity. The change in the body weight from baseline to 24 weeks is similar between the 2 treatment groups (-0.5 kg PBO+MET; -0.4 kg LI+MET). Conclusion, linagliptin 5 mg qd as add-on therapy in patients with T2DM inadequately controlled on metformin is well tolerated and produces significant and clinically meaningful improvements in glycemic control (reductions in HbA1c, FPG and 2h PPG without weight gain). Linagliptin as add-on therapy to metformin in patients with T2DM and insufficient glycemic control is well tolerated with the incidence of adverse events

Example 13

Linagliptin Improves Glycemic Control in Type 2 Diabetes Patients Inadequately Controlled by Metformin and Sulfonylurea without Weight Gain or Hypoglycemia

A multi-center, 24-week, randomized, double-blind, platrough levels in patients with mild and moderate renal 25 cebo-controlled, parallel group study examines the efficacy and safety of the DPP-4 inhibitor linagliptin (LI; 5 mg qd) in type 2 diabetes (T2DM) patients (pts) with insufficient glycemic control (HbA1c 7.0-10.0%) on the combination of metformin (MET) plus a sulfonylurea (SU). Effects of LI as 30 add-on are compared with placebo (PBO). All pts have a 2-wk PBO run-in before being randomized to LI+MET+SU (n=793) or PBO+MET+SU (n=265). Mean baseline characteristics are: HbA1c, 8.14% (SD 0.8); fasting plasma glucose (FPG), 160.1 mg/dL (36.6); age, 58.1 yrs (9.8); 35 BMI, 28.3 kg/m2 (4.7). Most of the pts (73.3%) have T2DM for >5 years before enrollment. The primary endpoint is the change from baseline in HbA1c after 24 weeks of treatment, adjusted for baseline HbA1c. After 24 weeks of treatment, the mean HbA1c for LI+MET+SU is -0.62% lower (p<0.0001) relative to PBO+MET+SU. The maximum mean HbA1c reduction with LI+MET+SU is seen at week 12 (-0.84%). Patients with baseline HbA1c ≥7.0% are >5-fold more likely to achieve a target HbA1c of <7.0% when treated with LI+MET+SU (29.2%) compared with PBO+ MET+SU (8.1%, odds ratio 5.5, p<0.0001) at 24 weeks. For the change in FPG, a statistically significant (p<0.0001) adjusted mean difference of -12.7 mg/dL is observed between LI+MET+SU and BPBO+MET+SU from baseline at week 24. Measures relating to β -cell function (fasting plasma insulin and HOMA-% B) along with HOMA-IR are significantly (p≤0.05) improved with LI+MET+SU compared with PBO+MET+SU. The proportion of patients that reported a severe adverse event (AE) is low for both LI+MET+SU and PBO+MET+SU groups (2.4% vs. 1.5%, the adjusted mean treatment difference between LI+MET 55 respectively). The most frequent AE reported more commonly in the LI+MET+SU group than in the PBO+MET+ SU group is hypoglycemia (22.7% vs. 14.8%, respectively). This is expected due to the combination with SU. No significant changes in weight are noted for either treatment group. Conclusion: Therapy with linagliptin added to the combination of metformin and a sulfonylurea is efficacious and safe in producing significant and clinically meaningful improvements in glycemic control in T2DM patients. Linagliptin may provide an additional option prior to insulin therapy in many patients for whom glycemia is insufficiently controlled with metformin plus a sulfonylurea agent. Linagliptin is shown to have a favorable safety and tolerablility

profile. However, when linagliptin is added on pre-existing sulfonylurea therapy, hypoglycemia may occur.

Example 14

Efficacy and Safety of Initial Combination Therapy with Linagliptin and Pioglitazone in Patients with Inadequately Controlled Type 2 Diabetes

A multi-center, 24-week, randomized, double-blind, pla- 10 cebo-controlled, parallel group study investigates the efficacy and safety of initial combination therapy with the DPP-4 inhibitor linagliptin (LI) and pioglitazone (PIO). Patients (pts) with type 2 diabetes mellitus (T2DM) and insufficient glycemic control (HbA1c 7.5-11.0%) who are 15 drug naïve or previously treated with any oral antihyperglycemic drug (OAD), are randomized to receive 5 mg LI plus 30 mg PIO qd (n=259) or 30 mg PIO plus placebo (PBO) qd (n=130). Patients do not take any OAD for at least 6 weeks before randomization. Mean baseline characteristics 20 (HbA1c 8.6%; fasting plasma glucose [FPG] 190 mg/dL; age 57.5 yrs; BMI 29.0 kg/m2) are similar between the groups. The primary endpoint is the change from baseline in HbA1c after 24 weeks of treatment, adjusted for baseline HbA1c and prior antidiabetic medication. After 24 weeks of 25 treatment, the adjusted mean change in HbA1c for the patients in the LI+ PIO group (full analysis set, last observation carried forward) is -1.06% (standard error (SE) ±0.06). The difference in the adjusted mean HbA1c for the LI+ PIO group compared with PBO+PIO is -0.51% 30 (p<0.0001; 95% confidence interval (CI), -0.71, -0.30). Reductions in FPG are also significantly greater for the LI+ PIO group compared with PBO+PIO with a treatment difference of -14.2 mg/dL (p<0.0001; 95% confidence interval (CI), -21.1, -7.3) at 24 weeks. Patients in the LI+ PIO group 35 are more likely to achieve a target HbA1c of <7% vs. those on PBO+PIO (42.9% vs. 30.5%, respectively, odds ratio 2.1; p=0.0051), as well as a reduction in HbA1c of \geq 0.5% (75%) vs. 50.8%, respectively, odds ratio 3.8; p<0.001). The proportion of patients that experienced at least one adverse 40 event (AE) is similar for both LI+ PIO and PBO+PIO groups (136, 52.5% vs. 53.1%, respectively). Hypoglycemia is rare, occurring in 3 patients (1.2%) in the LI+ PIO group and none in the PBO+PIO group. All hypoglycemic events are of mild intensity.

Conclusion: Initial combination therapy with linagliptin and pioglitazone shows significant and clinically meaningful improvements in FPG and HbA1c levels compared with PIO alone, along with a greater improvement in beta-cell function. Co-administration of linagliptin with pioglitazone is shown to be safe and well tolerated. Combination therapy with linagliptin and pioglitazone may provide an important synergistic initial treatment option for T2DM patients with inadequate glycemic control or those with renal impairment for whom metformin is contraindicated.

Example 15

Linagliptin Monotherapy Improves Glycemic Control in Japanese Patients with Type 2 Diabetes Mellitus Over 12 Weeks

A multi-center, 12-week, randomized, double-blind, placebo-controlled, parallel group study investigates the efficacy and safety of the DPP-4 inhibitor linagliptin (LI). 65 Effects of LI monotherapy (5 mg qd and 10 mg qd) are compared to placebo (PBO) in drug naïve or previously

64

treated Japanese patients (pts) with type 2 diabetes mellitus (T2DM) (baseline HbA1c 7.0-10.0%, if drug naïve; 7.0-9.0%, if previously treated). Before being randomized to LI 5 (n=159) or 10 mg (n=160), or PBO (n=80), all patients have a 2-week PBO run-in (patients on an oral antihyperglycemic drug have no medication for 2 weeks prior to run-in). Mean [SD] baseline characteristics and demographics (HbA1c, 8.0% [0.68]; fasting plasma glucose (FPG), 163.5 mg/dL [32.4]; BMI, 24.97 kg/m2 [3.86]; age, 60.0 yrs [9.7]) are similar in all groups. The primary endpoint is the change from baseline in HbA1c after 12 weeks. The differences of adjusted mean changes from baseline in HbA1c at week 12 are -0.87% for LI 5 mg vs. PBO (p<0.0001) and -0.88% for LI 10 mg vs. PBO (p<0.0001). Proportions of patients achieving HbA1c<7.0% after 12 wks are 26.4% for LI 5 mg and 35.7% for LI 10 mg vs. 10.0% for PBO. Proportions of patients whose HbA1c levels lower by at least 0.5% are 57.2% with LI 5 mg, 59.9% with LI 10 mg, and 8.8% with PBO. Both LI 5 mg and 10 mg show statistically significant difference compared with PBO (p<0.0001). FPG is significantly improved with both LI 5 and 10 mg compared to PBO: after 12 weeks, the differences of adjusted mean changes from baseline are -19.7 mg/dL for LI 5 mg vs. PBO (p<0.0001) and -20.4 mg/dL for LI 10 mg vs. PBO (p<0.0001). As indicated by changes in the proinsulin/insulin ratio (LI 5 mg, p=0.0065; LI 10 mg, p=0.0004), LI also significantly improves insulin secretion. The proportion of patients experiencing at least one adverse event (AE) is comparable among the three groups (56.0% LI 5 mg, 53.1% LI 10 mg and 56.3% PBO). Of those; 9.4%, 8.8% and 10.0%, respectively, are assessed as being drug-related. There are no investigator-defined hypoglycemic episodes. Body weight is unchanged with both LI 5 mg and 10 mg, -0.39 and -0.06 kg, respectively, which is not sigificantly different vs. PBO (-0.04 kg).

Conclusion: Linagliptin demonstrates a significant and clinically meaningful improvement in glycemic control, reflected in changes in HbA1c and FPG in Japanese patients with T2DM. Both linagliptin 5 and 10 mg doses have similar efficacy in lowering HbA1c and are well tolerated within this population. 5 mg linagliptin is the therapeutic dose in Japanese patients, which is identical to the therapeutic dose in Caucasians.

Example 16

Linagliptin Provides Superior Glycemic Control Compared to Voglibose as Monotherapy in Japanese Patients with Type 2 Diabetes

A multi-center, 26-week, randomized, double-blind, active-controlled, parallel group Study compares the efficacy and safety of the DPP-4 inhibitor linagliptin (LI) vs. the α-glucosidase inhibitor voglibose (VB) in drug naïve or previously treated Japanese patients (pts) with Type 2 diabetes mellitus (T2DM) (baseline HbA1c 7.0-10.0% if drug naïve, 7.0-9.0% if previously treated with an oral antihyperglycemic drug (OAD)).

Following a 2-week PBO run-in, patients are randomized to LI 5 (n=159) or 10 mg qd (n=160), or VB (0.2 mg tid; n=162). Any previous OAD treatment is stopped 2 weeks prior to run-in. Mean baseline [SD] characteristics and demographics (HbA1c, 8.01% [0.68]; fasting plasma glucose (FPG), 163.5 mg/dL [32.4]; BMI, 24.97 kg/m2 [3.86]; age, 60.0 yrs [9.7]) are similar across groups. The primary endpoint is the change from baseline in HbA1c after 26 weeks. The differences of adjusted mean changes from

baseline in HbA1c at week 26 are -0.32% for LI 5 mg vs. VB (p=0.0003) and -0.39% for LI 10 mg vs. VB (p<0.0001). Proportions of patients achieving HbA1c<7.0% after 26 weeks are 30.2% for LI 5 mg and 34.4% for LI 10 mg vs. 22.2% for VB. Proportions of patients whose HbA1c 5 level lowered by ≥0.5% are 57.2% and 53.5% for LI 5 and 10 mg, vs. 37.7% for VB. FPG is significantly improved with both LI 5 and 10 mg compared to VB: the differences of adjusted mean changes from baseline are -6.9 mg/dL for LI 5 mg vs. VB (p=0.02) and -9.8 mg/dL for LI 10 mg vs. 10 VB (p=0.0015). Both LI 5 mg and 10 mg show a significant decrease of HbA1c in patients previously treated with 1 OAD compared with VB (p=0.003 and p=0.0011, respectively). The occurrence of ≥1 adverse event (AE) is comparable between groups (72.3% LI 5 mg, 77.5% LI 10 mg 15 and 71.6% VB). Of the AEs, 11.3%, 10.6% and 18.5%, respectively, are assessed as drug related. Drug-related gastrointestinal disorders are more common in the VB (14.2%) than LI (8.2% 5 mg; 8.1% 10 mg) groups. In the VB group, 1 hypoglycemic episode is reported vs. none in the LI 20 groups.

Conclusion: Linagliptin monotherapy demonstrates greater efficacy than VB for improving glycemic control in Japanese patients with T2DM. Both linagliptin 5 mg and linagliptin 10 mg have comparable efficacy and show sta- 25 tistically significant decreases in HbA1c and FPG from baseline compared with VB after 26 weeks. Linagliptin is well tolerated in Japanese patients with T2DM compared to VB, with less gastrointestinal AEs, and may provide a valuable addition to the therapies available to this popula- 30 tion. 5 mg linagliptin is the therapeutic dose in Japanese patients, which is identical to the therapeutic dose in Caucasians.

Example 17

Linagliptin Restores β-Cell Function and Survival in Human Isolated Islets

Studies in diabetic animal models show that dipeptidyl 40 peptidase-4 (DPP-4) inhibitors reverse hyperglycemia and increase β-cell mass. Here, the role of linagliptin, a DPP-4 inhibitor on human β-cell function is investigated: Human isolated islets are exposed to increased glucose concentrations (5.5-33.3 mM), 0.5 mM palmitic acid, the mixture of 45 2 ng/mL IL-1 β or 1,000 U/mL IFN- γ for 4 days or 50 μM H2O2 for 8 h. Islets are pre-treated with 500 ng/mL Interleukin-1 Receptor Antagonist (IL-1Ra, which has been shown to restore β-cell function), 100 nM linagliptin or solvent for 1 h before exposure to the diabetic stimuli and 50 during the whole 4-day treatment period. At control conditions, islets secrete 3.8-fold more insulin at 16.7 mM than at 2.8 mM glucose. In contrast, stimulatory index is 1.9- and 2.4-fold decreased when islets are exposed to 11.1 mM and 33.3 mM glucose (P<0.05). Exposure of the islets to palmi- 55 affect the pharmacokinetics of linagliptin whereas it tate, cytokine mixture or H2O2 resulted in a 2.1-, 2.2- and 1.9-fold reduction of glucose stimulated insulin secretion (GSIS), respectively (P<0.05). Linagliptin significantly restores β-cell function at all conditions (1.9-, 2.5-, 3.3-, 1.9and 3.7-fold increase in GSIS at 11.1 or 33.3 mM glucose, 60 palmitic acid, cytokines or H2O2, P<0.05). IL-1Ra is similarly effective in restoring β-cell function at conditions of high glucose, palmitic acid and cytokines, but IL-1Ra failed to restore β-cell function at oxidative stress conditions induced by H2O2 treatment. Since loss of function is 65 mediated by oxidative stress, the nitrotyrosine concentration is measured in islet lysates. Nitrotyrosine levels are highly

66

elevated in human islets under all diabetic conditions (13-, 14-, 6-, 14- and 8-fold increased at 11.1 or 33.3 mM glucose, palmitic acid, cytokines or H2O2, P<0.05), while no elevated nitrotyrosine production is observed in islets treated with linagliptin.

In summary, it is shown that the DPP-4 inhibitor linagliptin has comparable protective effects on gluco-, lipo- and cytokinetoxicity as IL-1Ra and, in addition, could improve β-cell function under oxidative stress conditions and blocks apoptosis (induced by H2O2 treatment). The study provides evidence of a direct protective effect of linagliptin on β -cell survival and insulin secretion.

Example 18

Chronic Renal Disease does not Change the Pharmacokinetics of Linagliptin but Increases Exposure of Sitagliptin and Alogliptin in Rats

Renal impairment is a frequent complication of T2DM. The effect of chronic renal disease on the pharmacokinetics of dipeptidyl peptidase-4 inhibitors (linagliptin, sitagliptin, alogliptin) in a rat model of chronic renal insufficiency (5/6 nephrectomy, 5/6N) is investigated: Eight weeks after surgery rats are treated orally with inhibitors for 4 days. 5/6N causes a highly significantly (P<0.001) decrease of glomerular filtration rate measured by creatinin clearance (sham: 2510±210 mL/24 h; 5/6N: 1665±104.3 mL/24 h) and increases cystatin C levels (sham: 700±35.7 ng/mL; 5/6N: 1434±77.6 ng/mL). Tubular function is significantly (P<0.001) impaired as evidenced by plasma neutrophil gelatinase-associated lipocalin (NGAL), (sham: 286±23 ng/ml; 5/6N: 680±56.3 ng/ml) and β2 microglobulin (sham: 20.4±2.4 μg/mL; 5/6N, 33.3±1.34 μg/mL). DPP-4 activity is 35 comparable among groups.

Administration of linagliptin (0.5 and 7 µmol/kg) to 5/6N rats shows no significant change in AUC (0-∞): sham: 316±54.7 nmol*h/L; 5/6N: 257±21.54 nmol*h/L; P=0.771 and sham: 1252±372 nmol*h/L; 5/6N: 748±74.5 nmol*h/L; P=0.284, respectively. In contrast, both sitagliptin and alogliptin (7 µmol/kg) have significantly (P=0.0001 and P=0.039) higher (41% and 28%) AUC (0-∞): sitagliptin sham: 3690±103 nmol*h/L; 5/6N: 6238±423 nmol*h/L and alogliptin sham: 1772±225 nmol*h/L; 5/6N: 2445±166 nmol*h/L). Furthermore, no correlation of markers of tubular and glomerular functions with linagliptin AUC is observed. In contrast, sitagliptin significantly correlate with creatinin clearance (r2=0.374, P<0.05), cystatin C (r2=0.499, P<0.01), NGAL (r2=0.604, P<0.01) and $\beta 2$ microglobulin (r2=0.543, P<0.01). Alogliptin correlates less significantly with cystatin C (r2=0.376, P<0.05) and β2 microglobulin (r2=0.391, P<0.05) but not with creatinin clearance and NGAL.

These results demonstrate that renal impairment does not increases the exposure of sitagliptin and alogliptin. Therefore, in contrast to sitagliptin and alogliptin, linagliptin may not have to be dose-adjusted in patients with T2DM and renal impairment or diabetic nephopathy.

Further, linagliptin significantly inhibits mRNA expression of profibrotic factors, such as TGF-β1, T1MP-1 and collagen (Col3alpha1) in the heart of uremic rats, which factors are tissue fibrosis markers of cardiac fibrosis and are increased in uremic heart. Characteristic cardiomyopathy with intestinal expansion and fibrosis develops often in uremia. Thus, these antifibrotic properties of DPP-4 inhibitors may be used for the treatment of cardiac and renal

injury, uremic heart, cardiac fibrosis and/or cardiomyopathy with intestinal expansion and fibrosis associated with uremia in patients with type 2 diabetes. The antifibrotic action of linagliptin can be an additional benefit for patients with chronic kidney and/or heart diseases that often accompany 5 type 2 diabetes.

Example 19

Linagliptin Improves Hepatic Steatosis in Rodent Models

Hepatic steatosis is a hallmark of patients with Type 2 diabetes and non-alcoholic fatty liver disease (NAFLD). Linagliptin is a selective and non-renal excreted inhibitor of 15 dipeptidyl peptidase-4 (DPP-4).

In a model of diet-induced obesity (D10, fed for 2 and 3 months), the effect of 4 weeks therapy with linagliptin (3 and 30 mg/kg/day, n=10) is investigated. Liver lipid content is detected by magnetic resonance spectroscopy (MRS) in vivo 20 and by analysis of liver triglycerides ex vivo. Linagliptin inhibits DPP-4 activity significantly (P<0.001) by 67% to 80% and 79% to 89% (3 and 30 mg/kg/day, respectively) compared to controls. Blood glucose levels following an OGTT (AUC) are significantly (P<0.01) decreased ranging 25 from 16% to 20% (3 mg/kg/day) and 20% to 26% (30 mg/kg/day). Likewise, liver fat content (MRS detection) is significantly reduced. Changes in liver fat content are visible as early as 2 weeks on treatment. The correlation between liver lipid content as measured by MRS and hepatic triglyc- 30 eride levels as measured ex vivo is r2=0.565 (P<0.0001).

Furthermore, ob/ob mice are analyzed after 14 days of linagliptin treatment (3 mg/kg/day or control) and blinded histological scoring is performed (severity and grade of fat content, markers of inflammation). DPP-4 activity is inhib- 35 ited by 80% and blood glucose AUC reduction is 25% (P<0.05). The histological score reveals less hepatic steatosis and inflammation in the linagliptin group (2.2±0.13, n=9, P<0.01) versus control (3±0.18, n=10).

In conclusion, linagliptin significantly reduces liver fat 40 content and histological NAFLD in a high fat diet model. Linagliptin reverses liver triglyceride content and hepatic steatosis (with greater therapeutic impact when hepatic steatosis is more pronounced), The reversal of hepatic steatosis supports the use of linagliptin in patients with Type 45 2 diabetes as well as liver-associated diseases (NAFLD).

Example 20

Linagliptin Functionally Counteracts a Dysregulation in DPP-4 Expression In Diabetes-Impaired Wounds

Impaired wound healing is a major complication of diabetes mellitus. The dipeptidyl peptidase-4 (DPP-4) inhibitor 55 Samples linagliptin improves wound healing (as shown in ob/ob mice). The impact of linagliptin on inflammatory markers in wounded skin is examined and a rationale for the beneficial action of linagliptin on wound healing is provided:

Wounds of linagliptin (3 mg/kg/day) and mock-treated 60 ob/ob mice for the inflammatory markers COX-2 and MIP by RNase protection assays are investigated with no significant differences. Furthermore, linagliptin does not increase the number of apoptotic infiltrating F4/80-positive macrophages. Therefore, the expression of DPP-4 in the skin of 65 diabetic and non-diabetic animals is assessed. Immunohistochemistry (IHC) and immunoblots reveal a strong expres68

sion of DPP-4 in skin from healthy and diabetic (ob/ob) mice and keratinocytes as the major cellular source of the enzyme. In line, the localization of DPP-4 protein in the skin nicely correlates with whole body autoradiography obtained after [3H]-labelled linagliptin treatment. Analyzing DPP-4 expression in mice upon full-thickness excisional wounding it is found that in healthy mice, DPP-4 protein expression declines over 3 days after injury and the enzyme remains absent in the late phase of repair. Interestingly, skin injury leads to a strong down-regulation of DPP-4 expression in proliferating wound margin keratinocytes (IHC). In contrast, in acute wounds of diabetic mice any DPP-4 expression can not be observed. DPP-4 protein, however, is expressed in the late phase of wound repair. The inverse regulation of DPP-4 protein in diabetic versus non-diabetic skin provides a functional basis of the positive action of linagliptin in wound healing processes. Thus, improvement of the wound healing process mediated by a suitable DPP-4 inhibitor, such as linagliptin, depends on the compensation (inhibition) of a dysregulated DPP-4 in diabetic wounds rather than on the anti-glycemic or immunomodulatory effects thereof. Thus, a DPP-4 inhibitor being suitable for improving wound healing is such a DPP-4 inhibitor which can effectively bind to DPP-4 in the skin, e.g. to dysregulated DPP-4 in diabetic wounds, preferably in its therapeutic dose level.

Furthermore in this context, a DPP-4 inhibitor being suitable for improving wound healing, particularly in a type 2 diabetes patient, is such a DPP-4 inhibitor which can be applied topically to wounds, e.g. comprised in wound dressings or patches or creams or ointments. Thus, the present invention further provides topical devices for wounds, such as e.g. wound dressings or patches, comprising linagliptin and, optionally, one or more pharmaceutically acceptable carriers and/or excipients.

Example 21

Association Study (Genotyping TCF7L2, Treatment Response)

The polymorphisms and variants of the gene TCF7L2 as depicted in the Table i can be analyzed as described in the following procedure:

TABLE i

Gene, variant nucleotides and rs numbers.		
Gene	variant nucleotide	rs number
TCF7L2	c.382-41435 C > T c.483+9017 G > T, c.382-22060 A > G c.1102 C > G	rs7903146 rs12255372 rs10885406 rs731788

50

Patients' DNA samples (conc.: 50 ng/µl) in 96-well-plates are used for the analytical methods applied.

Genotyping by Direct Sanger Sequencing

Using gDNA as a template, locus specific DNA fragments are amplified by polymerase chain reaction (PCR).

PCR is carried out using an ABI BioRad® Tetrad PCR System. Quality of the PCR products is analyzed by agarose gel electrophoresis The purified PCR-products are used as templates in sequencing reactions According to the chain terminating methodology of Sanger et al. (1977), the analysis of DNA sequence is based on the termination of a growing DNA strand due to incorporation of a dye-labeled

2',3'-Dideoxyribonucleotidetriphosphate (ddNTP) by the DNA polymerase. Purified sequencing products are analyzed using an ABI PRISM® 3730 Genetic Analyzer.

Sequencing data are generated using the original ABI Software. The subsequent KB-basecalling as well as the assembly is performed using the Staden Software Package. KB-basecalling assigns quality values to all called bases of automated sequencer traces using KB-basecaller error probabilities. These quality values are used during assembling the single reads and are the basic requirement for calculating the sequence accuracy (Applied Biosystems, 3730/3730xl/ DNA Analyzer Sequencing Analysis Software Training).

A quality value (q) of 20 corresponds to an error probability (ep) of 1/100, a value of 30 to an ep of 1/1000 and so on. In the assembly phase those values are set against each other. In general sequencing is continued until each consensus base has a quality value (q) of 50 or more. This corresponds to an error probability (ep) of 1/100000. Due to the fact that most of the consensus bases have an even higher quality score than the minimal one, the calculated cumulative error probability for the finished sequence is again significantly lower. Sequencing data are uploaded and analyzed using the software segpatient from jsi-medical systems (version Seq Pilot 3.3.2, JSI medical systems GmbH, Friedhofstr. 5, 77971 Kippenheim, Germany).

Only traces that fulfill internal quality aspects are processed for further genotype analyses. Genotyping is carried out through the analysis of single polymorphisms rather than the analysis of the entire gene. Therefore genotyping results refer only to the variant positions depicted in Table i. Genotyping by TaqMan PCR

The TaqMan® technology comprises amplification of a PCR fragment with simultaneous detection of the degradation of a labelled probe. Probes are labelled at both ends with an allele-specific dye and a quencher. During the amplification reaction, the specifically hybridized probe is displaced by the DNA polymerase. This displacement occurs either as degradation through the 5' exonuclease activity of the polymerase in the case of a perfect match with the probe, or without degradation in the case of a mismatch. Upon degradation, the quencher and dye are separated and the fluorescence signal increased. An increase in the fluorescence signal is indicative for the presence of the respective allele. Fluorescence signals are recorded with the ABI PRISM 7700 system (Applied Biosystems).

In detail, a master mix is prepared containing all components for PCR reaction and aliquoted in the appropriate number of wells. Subsequently, DNA is added to each well according to the plate layout; except for no-template control

AB assay ID (rs7903146) C_29347861_10 SNP Context Sequence:

TAGAGAGCTAAGCACTTTTTAGATA[C/T]TATATAATTTAATTGCCGT

ATGAGG (SEO ID NO: 1)

The mastermix per sample contains:

Nuclease-free water	0.25 µl
2x PCR MasterMix	2.5 µl
20x Primer/Probe Mix	0.25 μl
DNA [10 ng/µl]	2 μl
In total:	5 µl

70

The cycling conditions are:

95° C. 95° C. 60° C.	10 min. 15 sec. 90 sec.	}	50 cycles	
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The TaqMan® pre- and post-reads of the AD are performed on the TaqMan® 7900HT Fast Real System. The SDS software V2.3 calculates the fluorescence measurements made during the plate read and plots Rn values based on the signals from each well. Using the software, it is determined which SNP alleles are present in each sample. NTC should be given as not determined.

Statistical Analyses

To assess the homogeneity of the treatment effect on the change from baseline of HbA1c after 24 weeks in the genotype subgroups defined by TCF7L2 SNP rs7903146 genotypes an analysis of covariance (ANCOVA) model including the treatment interaction with the covariate genotype is applied for pooled data over four studies. The statistical model includes 'Treatment', 'Genotype', 'Study', 'Wash-Out-Period for prior oral antidiabetic drugs (yes/no)', 'Race'. as well as the interaction 'Treatment*Genotype' as fixed effects and 'HbA1c baseline' as a linear covariate. The ANCOVA model provides estimates for the mean change from baseline in HbA1c after 24 weeks of therapy for the different genotypes taking baseline clinical and demographic information into account.

Model based pair-wise comparisons between wild-type homozygous (genotype CC) and heterozygous (genotype CT) or rare homozygous (genotype TT) individuals on linagliptin or combination treatment (linagliptin+pioglitazone, linagliptin+metformin, linagliptin+metformin+a sulphonylurea) are performed.

Additionally the results of the corresponding ANCOVA models without 'Genotype' and 'Treatment*Genotype' fixed effects are given for the whole patient population of the studies (full analysis set, FAS) as well as for the subpopulation for which genetic analyses are performed (full analysis set for pharmacogenetic analyses, FASG) to demonstrate comparability of the observed effects.

The statistical evaluation is prepared using the software packages SAS Version 9.2 (SAS Institute Inc., Cary, N.C., 45 USA) and S-PLUS® 8.0 (Insightful Corp., Seattle, Wash., USA).

FIG. 1 shows mean values and 95% confidence intervals for baseline HbA1c values for the whole patient population of the studies (full analysis set, FAS), for the subpopulation for which genetic analyses are performed (full analysis set for pharmacogenetic analyses, FASG), as well as for the subgroups defined by genotype (CC, CT, TT) of this subpopulation. The numbers of patients for placebo control and linagliptin treatment are given in braces.

FIG. 2 shows a statistical association between TCF7L2 SNP rs7903146 genotypes with a likelihood of a favorable response in CC/CT genotype carriers to the administration of a therapeutically-effective amount of linagliptin or linagliptin in combination with other oral antidiabetic therapy.

Results are shown as point estimates and 95% confidence intervals for the mean change in HbA1c from baseline [%] after 24 weeks as estimated by ANCOVA models. The results are given for the whole patient population of the studies (full analysis set, FAS), for the subpopulation for which genetic analyses are performed (full analysis set for pharmacogenetic analyses, FASG), as well as for the subgroups defined by genotype (CC, CT, TT) of this subpopu-

lation. The numbers of patients for placebo control and linagliptin treatment are given in braces.

Point estimates and 95% confidence intervals for the differences in changes in HbA1c from baseline [%] for the comparison of between wild-type homozygous (genotype CC) and heterozygous (genotype CT) or rare homozygous (genotype TT) individuals on linagliptin treatment or combination treatment (linagliptin+pioglitazone, linagliptin+metformin, linagliptin+metformin+a sulphonylurea) are shown as well. They result in a statistically significant difference between TT and CC (p value=0.0192). (Other pairwise comparisons: CT vs. CC: p=0.4359; CT vs. TT: p=0.0712).

This indicates a significant association between the wildtype homozygous genotype and lower HbA1c on treatment.

FIG. 1 Baseline HbA1c values for the whole patient population of the studies (full analysis set, FAS), for the subpopulation for which genetic analyses are performed (full analysis set for pharmacogenetic analyses, FASG), as well as for the subgroups defined by SNP rs7903146 in TCF7L2 genotypes (CC, CT, TT) of this subpopulation.

FIG. 2 Association of SNP rs7903146 in TCF7L2 with linagliptin response

EXAMPLES OF FORMULATIONS

The following examples of formulations, which may be obtained analogously to methods known in the art, serve to illustrate the present invention more fully without restricting it to the contents of these examples. The term "active substance" denotes one or more compounds according to the invention, i.e. denotes a DPP-4 inhibitor or a second or third antidiabetic compound according to this invention or a combination of two or three of said active ingredients, for example selected from the combinations as listed in the Table 1 or 2. Additional suitable formulations for the DPP-4 inhibitor linagliptin may be those formulations disclosed in the application WO 2007/128724, the disclosure of which is incorporated herein in its entirety. Additional suitable formulations for the other DPP-4 inhibitors may be those formulations which are available on the market, or formulations described in the patent applications cited above in paragraph "background of the invention", or those described in the literature, for example as disclosed in current issues of "Rote Liste®" (Germany) or of "Physician's Desk Reference".

Example 1

Dry Ampoule Containing 75 Mg of Active Substance Per 10 ml

Composition:

Active substance	75.0 mg
Mannitol	50.0 mg
water for injections	ad 10.0 ml

Preparation:

Active substance and mannitol are dissolved in water. After packaging the solution is freeze-dried. To produce the 65 solution ready for use, the product is dissolved in water for injections.

72

Example 2

Dry Ampoule Containing 35 Mg of Active Substance Per 2 ml

Composition:

	Active substance	35.0 mg	
0	Mannitol	100.0 mg	
	water for injections	ad 2.0 ml	

Preparation:

Active substance and mannitol are dissolved in water. ⁵ After packaging, the solution is freeze-dried.

To produce the solution ready for use, the product is dissolved in water for injections.

Example 3

Tablet Containing 50 mg of Active Substance

Composition:

25

50

60

(1) Active substance (2) Mannitol (3) Maize starch (4) Polyvinylpyrrolidone (5) Magnesium stearate	50.0 mg 98.0 mg 50.0 mg 15.0 mg 2.0 mg
_	215.0 mg

Preparation:

(1), (2) and (3) are mixed together and granulated with an aqueous solution of (4). (5) is added to the dried granulated material. From this mixture tablets are pressed, biplanar, faceted on both sides and with a dividing notch on one side. Diameter of the tablets: 9 mm.

Example 4

Tablet Containing 350 mg of Active Substance Preparation:

350.0 mg
136.0 mg
80.0 mg
30.0 mg
4.0 mg
600.0 mg

(1), (2) and (3) are mixed together and granulated with an aqueous solution of (4). (5) is added to the dried granulated
 material. From this mixture tablets are pressed, biplanar, faceted on both sides and with a dividing notch on one side. Diameter of the tablets: 12 mm.

Example 5

Capsules Containing 50 Mg of Active Substance

Composition:

(1) Active substance(2) Dried maize starch	50.0 mg 58.0 mg

(3) Mannitol	50.0 mg
(4) Magnesium stearate	2.0 mg
	160.0 mg

Preparation:

(1) is triturated with (3). This trituration is added to the mixture of (2) and (4) with vigorous mixing. This powder mixture is packed into size 3 hard gelatin capsules in a capsule filling machine.

Example 6

Capsules Containing 350 mg of Active Substance

Composition:

(1) Active substance	350.0 mg	20
(2) Dried maize stard	ch 46.0 mg	
(3) Mannitol	30.0 mg	
(4) Magnesium stear	ate 4.0 mg	
	430.0 mg	25

Preparation:

(1) is triturated with (3). This trituration is added to the mixture of (2) and (4) with vigorous mixing. This powder mixture is packed into size 0 hard gelatin capsules in a capsule filling machine.

74

- GLP-1 or GLP-1 analogues, and insulin or insulin analogues, optionally in combination,
- wherein the patient has insufficient glycemic control despite diet and exercise or despite previous monotherapy with the second or the third antidiabetic agent, or
- wherein the patient has insufficient glycemic control despite diet and exercise or despite previous dual therapy with the second and the third antidiabetic agent.
- 2. The method according to claim 1 wherein the patient is an individual diagnosed of one or more of the conditions selected from the group consisting of overweight, obesity, visceral obesity and abdominal obesity.
- 3. The method according to claim 1 wherein the patient is an individual who shows one, two or more of the following conditions:
 - (a) a fasting blood glucose or serum glucose concentration greater than 110 mg/dL, in particular greater than 125 mg/dL;
 - (b) a postprandial plasma glucose equal to or greater than 140 mg/dL;
 - (c) an HbA1c value equal to or greater than 6.5%, in particular equal to or greater than 7.0%.
- **4**. The method according to claim **1** wherein the patient is an individual wherein one, two, three or more of the following conditions are present:
 - (a) obesity, visceral obesity and/or abdominal obesity,
 - (b) triglyceride blood level ≥150 mg/dL,
 - (c) HDL-cholesterol blood level <40 mg/dL in female patients and <50 mg/dL in male patients,</p>

SEQUENCE LISTING

<160> NUMBER OF SEQ ID NOS: 1

<210> SEQ ID NO 1

<211> LENGTH: 51 <212> TYPE: DNA

<213> ORGANISM: Homo sapiens

<400> SEQUENCE: 1

tagagageta ageaettttt agataytata taatttaatt geegtatgag g

51

The invention claimed is:

- 1. A method for treating type 2 diabetes mellitus in a patient in need thereof, wherein said patient is responsive to treatment of type 2 diabetes and has at least one variation in the TCF7L2 gene, wherein said at least one variation in the TCF7L2 gene is a single nucleotide polymorphism (SNP) in the gene coding for TCF7L2, wherein the SNP is selected from the group consisting of rs7903146, rs12255372 and rs10885406, said method comprising
 - testing to determine if the patient has said at least one variation in the TCF7L2 gene, and
 - administering to the patient having the at least one variation in the TCF7L2 gene a DPP-4 inhibitor selected from linagliptin, sitagliptin, vildagliptin, alogliptin, saxagliptin, teneligliptin and dutogliptin; a second antidiabetic agent selected from biguanides, thiazolidindiones, sulfonylureas, glinides, inhibitors of alphaglucosidase, GLP-1 or GLP-1 analogues, and insulin or insulin analogues: and, optionally, a third antidiabetic 65 agent selected from biguanides, thiazolidindiones, sulfonylureas, glinides, inhibitors of alpha-glucosidase,

- (d) a systolic blood pressure ≥130 mm Hg and a diastolic blood pressure ≥85 mm Hg,
- (e) a fasting blood glucose level ≥110 mg/dL.
- 5. The method according to claim 1 wherein the patient 50 has insufficient glycemic control despite diet and exercise or despite monotherapy with the second or the third antidiabetic agent.
 - 6. The method according to claim 1 wherein the patient has insufficient glycemic control despite diet and exercise or despite dual therapy with the second and the third antidiabetic agent.
 - 7. The method according to claim 1 wherein the patient has been diagnosed with at least one T allele of SNP rs7903146 of TCF7L2.
 - **8**. The method according to claim **1** wherein the patient has been diagnosed with two T alleles of SNP rs7903146 of TCF7L2.
 - **9**. The method according claim **1** wherein the DPP-4 inhibitor is linagliptin.
 - 10. The method according to claim 1 wherein the DPP-4 inhibitor is linagliptin, and the second antidiabetic agent is metformin or pioglitazone.

45

- 11. The method according to claim 1 wherein the patient suffers from or has been diagnosed with type 2 diabetes mellitus.
- 12. The method of claim 1, wherein said DPP-4 inhibitor is administered in combination with a third antidiabetic $_5$ agent.
- 13. A method of using a pharmaceutical composition comprising
 - (a) a DPP-4 inhibitor selected from linagliptin, sitagliptin, vildagliptin, alogliptin, saxagliptin, teneligliptin and dutogliptin,
 - (b) a second antidiabetic agent selected from the group G3 consisting of biguanides, thiazolidindiones, sulfonylureas, glinides, inhibitors of alpha-glucosidase, GLP-1 or GLP-1 analogues, and insulin or insulin analogues, and, optionally,
 - (c) a third antidiabetic agent being different from (b) selected from the group G3 consisting of biguanides, thiazolidindiones, sulfonylureas, glinides, inhibitors of alpha-glucosidase, GLP-1 or GLP-1 analogues, and insulin or insulin analogues,

or a pharmaceutically acceptable salt thereof,

for treating a type 2 diabetes in a patient in need thereof susceptible to and in need of such treatment, wherein said patient is responsive to treatment of type 2 diabetes, the method comprising administering said pharmaceutical composition to said patient, wherein

- the patient has been diagnosed with a variation in the TCF7L2 gene, wherein said variation in the TCF7L2 gene is a single nucleotide polymorphism (SNP) in the gene coding for TCF7L2 wherein said SNP is selected from the group consisting of rs7903146, rs12255372 and rs10885406
- wherein the patient has insufficient glycemic control despite diet and exercise or despite previous monotherapy with the second or the third antidiabetic agent, 35
- wherein the patient has insufficient glycemic control despite diet and exercise or despite previous dual therapy with the second and the third antidiabetic agent.

- 14. A method of using a pharmaceutical composition comprising
 - (a) DPP-4 inhibitor selected from linagliptin, sitagliptin, vildagliptin, alogliptin, saxagliptin, teneligliptin and dutogliptin,
 - (b) a second antidiabetic agent selected from the group G3 consisting of biguanides, thiazolidindiones, sulfonylureas, glinides, inhibitors of alpha-glucosidase, GLP-1 or GLP-1 analogues, and insulin or insulin analogues, and, optionally,
 - (c) a third antidiabetic agent being different from (b) selected from the group G3 consisting of biguanides, thiazolidindiones, sulfonylureas, glinides, inhibitors of alpha-glucosidase, GLP-1 or GLP-1 analogues, and insulin or insulin analogues,
 - or a pharmaceutically acceptable salt thereof,
 - for treating type 2 diabetes mellitus in a patient in need thereof, wherein said patient is responsive to treatment of type 2 diabetes, said method comprising
 - (i) testing to determine if the patient has at least one single nucleotide polymorphism (SNP) in the gene encoding for TCF7L2, wherein the at least one SNP is selected from rs7903146, rs12255372 and rs10885406

and.

- (ii) administering the pharmaceutical composition to the patient having the at least one SNP in the gene coding for TCF7L2.
- wherein the patient has insufficient glycemic control despite diet and exercise or despite previous monotherapy with the second or the third antidiabetic agent;

wherein the patient has insufficient glycemic control despite diet and exercise or despite previous dual therapy with the second and the third antidiabetic agent.

15. The method according to claim **14** wherein the patient has been diagnosed with TCF7L2 rs7903146 CT or TT risk genotype.

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